

Learning statistics with R:
A tutorial for psychology students and other beginners
(Version 0.3.1)

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This book was brought to you today by the letter ‘R’.

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Preface

There's a part of me that really doesn't want to publish this book. It's not finished.

And when I say that, I mean it. The referencing is spotty at best, the chapter summaries are just lists of section titles, there's no index, there are no exercises for the reader, the organisation is suboptimal, and the coverage of topics is just not comprehensive enough for my liking. Additionally, there are sections with content that I'm not happy with, figures that really need to be redrawn, and I've had almost no time to hunt down inconsistencies, typos, or errors. In other words, *this book is not finished*. If I didn't have a looming teaching deadline and a baby due in a few weeks, I really wouldn't be making this available at all.

What this means is that if you are an academic looking for teaching materials, a Ph.D. student looking to learn R, or just a member of the general public interested in statistics, I would advise you to be cautious. What you're looking at is a first draft, and it may not serve your purposes. If we were living in the days when publishing was expensive and the internet wasn't around, I would never consider releasing a book in this form. The thought of someone shelling out \$80 for this (which is what a commercial publisher told me it would retail for when they offered to distribute it) makes me feel more than a little uncomfortable. However, it's the 21st century, so I can post the pdf on my website for free, and I can distribute hard copies via a print-on-demand service for less than half what a textbook publisher would charge. And so my guilt is assuaged, and I'm willing to share! With that in mind, you can obtain free soft copies and cheap hard copies online, from the following webpages:

Soft copy: ua.edu.au/ccs/teaching/lsl

Hard copy: www.lulu.com/content/13570633

Even so, the warning still stands: what you are looking at is Version 0.3 of a work in progress. If and when it hits Version 1.0, I would be willing to stand behind the work and say, yes, this is a textbook that I would encourage other people to use. At that point, I'll probably start shamelessly flogging the thing on the internet and generally acting like a tool. But until that day comes, I'd like it to be made clear that I'm really ambivalent about the work as it stands.

All of the above being said, there is one group of people that I can enthusiastically endorse this book to: the psychology students taking our undergraduate research methods classes (DRIP and DRIP:A) in 2013. For you, this book is ideal, because it was written to accompany your stats lectures. If a problem arises due to a shortcoming of these notes, I can and will adapt content on the fly to fix that problem. Effectively, you've got a textbook written specifically for your classes, distributed for free (electronic copy) or at near-cost prices (hard copy). Better yet, the notes have been tested: Version 0.1 of these notes was used in the 2011 class, Version 0.2 was used in the 2012 class, and now you're looking at the new and improved Version 0.3. I'm not saying these notes are titanium plated awesomeness on a stick – though if *you* wanted to say so on the student evaluation forms, then you're totally welcome to – because they're not. But I am saying that they've been tried out in previous years and they seem to work okay. Besides, there's a group of us around to troubleshoot if any problems come up, and you can guarantee that at least *one* of your lecturers has read the whole thing cover to cover!

Okay, with all that out of the way, I should say something about what the book aims to be. At its core, it is an introductory statistics textbook pitched primarily at psychology students. As such, it covers the standard topics that you'd expect of such a book: study design, descriptive statistics, the theory of hypothesis testing, *t*-tests, χ^2 tests, ANOVA and regression. However, there are also several chapters devoted to the R statistical package, including a chapter on data manipulation and another one on scripts and programming. Moreover, when you look at the content presented in the book, you'll notice a lot of topics that are traditionally swept under the carpet when teaching statistics to psychology students. The Bayesian/frequentist divide is openly discussed in the probability chapter, and the disagreement between Neyman and Fisher about hypothesis testing makes an appearance. The difference between probability

and density is discussed. A detailed treatment of Type I, II and III sums of squares for unbalanced factorial ANOVA is provided. And if you have a look in the Epilogue, it should be clear that my intention is to add a lot more advanced content.

My reasons for pursuing this approach are pretty simple: the students can handle it, and they even seem to enjoy it. Over the last few years I've been pleasantly surprised at just how little difficulty I've had in getting undergraduate psych students to learn R. It's certainly not easy for them, and I've found I need to be a little charitable in setting marking standards, but they do eventually get there. Similarly, they don't seem to have a lot of problems tolerating ambiguity and complexity in presentation of statistical ideas, as long as they are assured that the assessment standards will be set in a fashion that is appropriate for them. So if the students can handle it, why *not* teach it? The potential gains are pretty enticing. If they learn R, the students get access to CRAN, which is perhaps the largest and most comprehensive library of statistical tools in existence. And if they learn about probability theory in detail, it's easier for them to switch from orthodox null hypothesis testing to Bayesian methods if they want to. Better yet, they learn data analysis skills that they can take to an employer without being dependent on expensive and proprietary software.

Sadly, this book isn't the silver bullet that makes all this possible. It's a work in progress, and maybe when it is finished it will be a useful tool. One among many, I would think. There are a number of other books that try to provide a basic introduction to statistics using R, and I'm not arrogant enough to believe that mine is better. Still, I rather like the book, and maybe other people will find it useful, incomplete though it is.

Dan Navarro
January 13, 2013

Part I.

Background

1. Why do we learn statistics?

*"Thou shalt not answer questionnaires
Or quizzes upon World Affairs,
Nor with compliance
Take any test. Thou shalt not sit
With statisticians nor commit
A social science"*

– W.H. Auden

1.1

On the psychology of statistics

To the surprise of many students, statistics is a fairly significant part of a psychological education. To the surprise of no-one, statistics is very rarely the *favourite* part of one's psychological education. After all, if you really loved the idea of doing statistics, you'd probably be enrolled in a statistics class right now, not a psychology class. So, not surprisingly, there's a pretty large proportion of the student base that isn't happy about the fact that psychology has so much statistics in it. In view of this, I thought that the right place to start might be to answer some of the more common questions that people have about stats...

A big part of this issue at hand relates to the very idea of statistics. What is it? What's it there for? And why are scientists so bloody obsessed with it? These are all good questions, when you think about it. So let's start with the last one. As a group, scientists seem to be bizarrely fixated on running statistical tests on everything. In fact, we use statistics so often that we sometimes forget to explain to people why we do. It's a kind of article of faith among scientists – and especially social scientists – that your findings can't be trusted until you've done some stats. Undergraduate students might be forgiven for thinking that we're all completely mad, because no-one takes the time to answer one very simple question:

Why do you do statistics? Why don't scientists just use common sense when evaluating evidence?

It's a naive question in some ways, but most good questions are. There's a lot of good answers to it,¹ but for my money, the best answer is a really simple one: we don't trust ourselves enough. We worry that

¹Including the suggestion that common sense is in short supply among scientists.

we're human, and susceptible to all of the biases, temptations and frailties that all humans suffer from. Much of statistics is basically a safeguard. Using "common sense" to evaluate evidence means trusting gut instincts, relying on verbal arguments and on using the raw power of human reason to come up with the right answer. Most scientists don't think this approach is likely to work.

In fact, come to think of it, this sounds a lot like a psychological question to me, and since I do work in a psychology department, it seems like a good idea to dig a little deeper here. Is it really plausible to think that this "common sense" approach is very trustworthy? Verbal arguments have to be constructed in language, and all languages have biases – some things are harder to say than others, and not necessarily because they're false (e.g., quantum electrodynamics is a good theory, but hard to explain in words). The instincts of our "gut" aren't designed to solve scientific problems, they're designed to handle day to day inferences – and given that biological evolution is slower than cultural change, we should say that they're designed to solve the day to day problems for a *different world* than the one we live in. Most fundamentally, reasoning sensibly requires people to engage in "induction", making wise guesses and going beyond the immediate evidence of the senses to make generalisations about the world. If you think that you can do that without being influenced by various distractors, well, I have a bridge in Brooklyn I'd like to sell you. Heck, as the next section shows, we can't even solve "deductive" problems (ones where no guessing is required) without being influenced by our pre-existing biases.

1.1.1 The curse of belief bias

People are mostly pretty smart. We're certainly smarter than the other species that we share the planet with (though many people might disagree). Our minds are quite amazing things, and we seem to be capable of the most incredible feats of thought and reason. That doesn't make us perfect though. And among the many things that psychologists have shown over the years is that we really do find it hard to be neutral, to evaluate evidence impartially and without being swayed by pre-existing biases. A good example of this is the **belief bias effect** in logical reasoning: if you ask people to decide whether a particular argument is logically valid (i.e., conclusion would be true if the premises were true), we tend to be influenced by the believability of the conclusion, even when we shouldn't. For instance, here's a valid argument where the conclusion is believable:

No cigarettes are inexpensive (Premise 1)
Some addictive things are inexpensive (Premise 2)
Therefore, some addictive things are not cigarettes (Conclusion)

And here's a valid argument where the conclusion is not believable:

No addictive things are inexpensive (Premise 1)
Some cigarettes are inexpensive (Premise 2)
Therefore, some cigarettes are not addictive (Conclusion)

The logical *structure* of argument #2 is identical to the structure of argument #1, and they're both valid. However, in the second argument, there are good reasons to think that premise 1 is incorrect, and as a result it's probably the case that the conclusion is also incorrect. But that's entirely irrelevant to the topic at hand: an argument is deductively valid if the conclusion is a logical consequence of the premises. That is, a valid argument doesn't have to involve true statements.

On the other hand, here's an invalid argument that has a believable conclusion:

No addictive things are inexpensive (Premise 1)
Some cigarettes are inexpensive (Premise 2)
Therefore, some addictive things are not cigarettes (Conclusion)

And finally, an invalid argument with an unbelievable conclusion:

No cigarettes are inexpensive (Premise 1)
Some addictive things are inexpensive (Premise 2)
Therefore, some cigarettes are not addictive (Conclusion)

Now, suppose that people really are perfectly able to set aside their pre-existing biases about what is true and what isn't, and purely evaluate an argument on its logical merits. We'd expect 100% of people to say that the valid arguments are valid, and 0% of people to say that the invalid arguments are valid. So if you ran an experiment looking at this, you'd expect to see data like this:

	conclusion feels true	conclusion feels false
argument is valid	100% say “valid”	100% say “valid”
argument is invalid	0% say “valid”	0% say “valid”

If the psychological data looked like this (or even a good approximation to this), we might feel safe in just trusting our gut instincts. That is, it'd be perfectly okay just to let scientists evaluate data based on their common sense, and not bother with all this murky statistics stuff. However, you guys have taken psych classes, and by now you probably know where this is going ...

In a classic study, [Evans, Barston, and Pollard \(1983\)](#) ran an experiment looking at exactly this. What they found is that when pre-existing biases (i.e., beliefs) were in agreement with the structure of the data, everything went the way you'd hope:

	conclusion feels true	conclusion feels false
argument is valid	92% say “valid”	
argument is invalid		8% say “valid”

Not perfect, but that's pretty good. But look what happens when our intuitive feelings about the truth of the conclusion run against the logical structure of the argument:

	conclusion feels true	conclusion feels false
argument is valid	92% say “valid”	46% say “valid”
argument is invalid	92% say “valid”	8% say “valid”

Oh dear, that's not as good. Apparently, when people are presented with a strong argument that contradicts our pre-existing beliefs, we find it pretty hard to even perceive it to be a strong argument (people only did so 46% of the time). Even worse, when people are presented with a weak argument that agrees with our pre-existing biases, almost no-one can see that the argument is weak (people got that one wrong 92% of the time!)²

If you think about it, it's not as if these data are horribly damning. Overall, people did do better than chance at compensating for their prior biases, since about 60% of people's judgements were correct (you'd expect 50% by chance). Even so, if you were a professional “evaluator of evidence”, and someone came along and offered you a magic tool that improves your chances of making the right decision from 60% to (say) 95%, you'd probably jump at it, right? Of course you would. Thankfully, we actually do have a tool that can do this. But it's not magic, it's statistics. So that's reason #1 why scientists love

²In my more cynical moments I feel like this fact alone explains 95% of what I read on the internet.

statistics. It's just *too easy* for us to "believe what we want to believe"; so if we want to "believe in the data" instead, we're going to need a bit of help to keep our personal biases under control. That's what statistics does: it helps keep us honest.

1.2

The cautionary tale of Simpson's paradox

The following is a true story. In 1973, the University of California, Berkeley got into a lot of trouble over its admissions of students into postgraduate courses. For those of you who don't know how the US university system works, it's very different to Australia. Unlike Australia, there's no centralisation, and university admission is based on many different factors besides grades. Moreover, each university makes its own decisions about who to send offers to, using its own decision process to do so. So, when the 1973 admissions data looked like this...

	Number of applicants	Percent admitted
Males	8442	44%
Females	4321	35%

...they got sued. Given that there were nearly 13,000 applicants, a difference of 9% in admission rates between males and females is just way too big to be a coincidence. Pretty compelling data, right? And if I were to say to you that these data *actually* reflect a weak bias in favour of females, you'd probably think that I was either crazy or sexist.

Oddly, it's actually sort of true ...after Berkeley got sued, people started looking *very* carefully at the admissions data (Bickel, Hammel, & O'Connell, 1975). And remarkably, when they looked at it on a department by department basis, it turned out that most of the departments actually had a slightly *higher* success rate for female applicants than for male applicants. The table below shows the admission figures for the six largest departments (with the names of the departments removed for privacy reasons):

Department	Males		Females	
	Applicants	Percent admitted	Applicants	Percent admitted
A	825	62%	108	82%
B	560	63%	25	68%
C	325	37%	593	34%
D	417	33%	375	35%
E	191	28%	393	24%
F	272	6%	341	7%

Remarkably, most departments had a *higher* rate of admissions for females than for males! Yet the overall rate of admission across the university for females was *lower* than for males. How can this be? How can both of these statements be true at the same time?

Here's what's going on. Firstly, notice that the departments are *not* equal to one another in terms of their admission percentages: some departments (e.g., engineering, chemistry) tended to admit a high percentage of the qualified applicants, whereas others (e.g., English) tended to reject most of the candidates, even if they were high quality. So, among the six departments shown above, notice that department A is the most generous, followed by B, C, D, E and F in that order. Next, notice that males and females tended to apply to different departments. If we rank the departments in terms of the total number of male applicants, we get **A>B>D>C>F>E** (the "easy" departments are in bold). On the whole, males tended to apply to the departments that had high admission rates. Now compare this to how the female

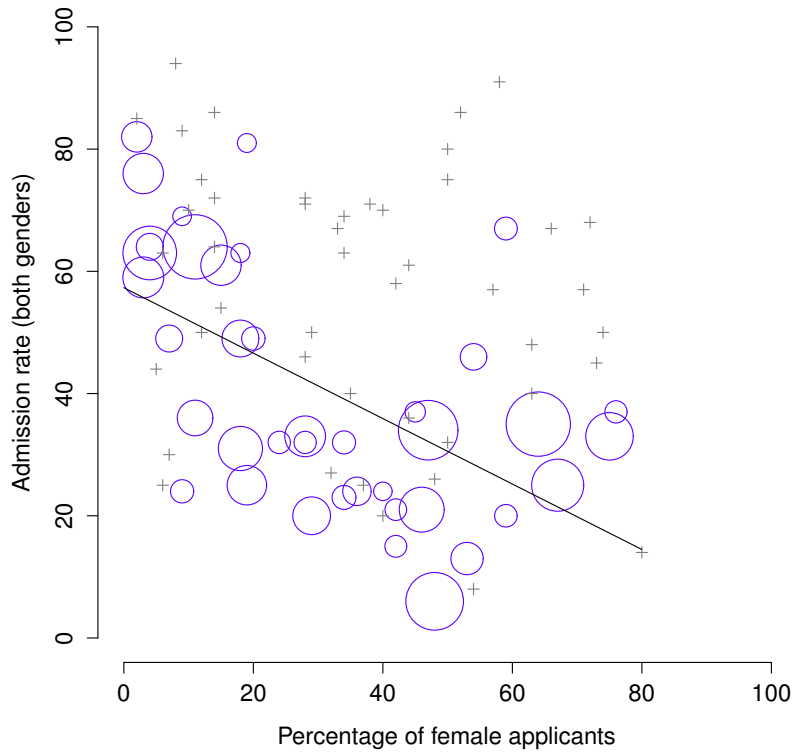


Figure 1.1: The Berkeley 1973 college admissions data. This figure plots the admission rate for the 85 departments that had at least one female applicant, as a function of the percentage of applicants that were female. The plot is a redrawing of Figure 1 from [Bickel et al. \(1975\)](#). Circles plot departments with more than 40 applicants; the area of the circle is proportional to the total number of applicants. The crosses plot department with fewer than 40 applicants.

applicants distributed themselves. Ranking the departments in terms of the total number of female applicants produces a quite different ordering C>E>D>F>**A**>**B**. In other words, what these data seem to be suggesting is that the female applicants tended to apply to “harder” departments. And in fact, if we look at all Figure 1.1 we see that this trend is systematic, and quite striking. This effect is known as **Simpson’s paradox**. It’s not common, but it does happen in real life, and most people are very surprised by it when they first encounter it, and many people refuse to even believe that it’s real. It is very real. And while there are lots of very subtle statistical lessons buried in there, I want to use it to make a much more important point . . . doing research is hard, and there are *lots* of subtle, counterintuitive traps lying in wait for the unwary. That’s reason #2 why scientists love statistics, and why we teach research methods. Because science is hard, and the truth is sometimes cunningly hidden in the nooks and crannies of complicated data.

Before leaving this topic entirely, I want to point out something else really critical that is often overlooked in a research methods class. Statistics only solves *part* of the problem. Remember that we started all this with the concern that Berkeley’s admissions processes might be unfairly biased against female

applicants. When we looked at the “aggregated” data, it did seem like the university was discriminating against women, but when we “disaggregate” and looked at the individual behaviour of all the departments, it turned out that the actual departments were, if anything, slightly biased in favour of women. The gender bias in admissions was entirely caused by the fact that women tended to self-select for harder departments. From a purely legal perspective, that puts the university in the clear. Postgraduate admissions are determined at the level of the individual department (and there are very good reasons to do that), and at the level of individual departments, the decisions are more or less unbiased (the weak bias in favour of females at that level is small, and not consistent across departments). Since the university can’t dictate which departments people choose to apply to, and the decision making takes place at the level of the department it can hardly be held accountable for any biases that those choices produce.

That was the basis for my somewhat glib remarks, but that’s not exactly the whole story, is it? After all, if we’re interested in this from a more sociological and psychological perspective, we might want to ask *why* there are such strong gender differences in applications. Why do males tend to apply to engineering more often than females, and why is this reversed for the English department? And why is it the case that the departments that tend to have a female-application bias tend to have lower overall admission rates than those departments that have a male-application bias? Might this not still reflect a gender bias, even though every single department is itself unbiased? It might. Suppose, hypothetically, that males preferred to apply to “hard sciences” and females prefer “humanities”. And suppose further that the reason for why the humanities departments have low admission rates is because the government doesn’t want to fund the humanities (Ph.D. places, for instance, are often tied to government funded research projects). Does that constitute a gender bias? Or just an unenlightened view of the value of the humanities? What if someone at a high level in the government cut the humanities funds because they felt that the humanities are “useless chick stuff”. That seems pretty blatantly gender biased. None of this falls within the purview of statistics, but it matters to the research project. If you’re interested in the overall structural effects of subtle gender biases, then you probably want to look at *both* the aggregated and disaggregated data. If you’re interested in the decision making process at Berkeley itself then you’re probably only interested in the disaggregated data.

In short there are a lot of critical questions that you can’t answer with statistics, but the answers to those questions will have a huge impact on how you analyse and interpret data. And this is the reason why you should always think of statistics as a *tool* to help you learn about your data, no more and no less. It’s a powerful tool to that end, but there’s no substitute for careful thought.

1.3

Statistics in psychology

I hope that the discussion above helped explain why science in general is so focused on statistics. But I’m guessing that you have a lot more questions about what role statistics plays in psychology, and specifically why psychology classes always devote so many lectures to stats. So here’s my attempt to answer a few of them...

• Why does psychology have so much statistics?

To be perfectly honest, there’s a few different reasons, some of which are better than others. The most important reason is that psychology is a statistical science. What I mean by that is that the “things” that we study are *people*. Real, complicated, gloriously messy, infuriatingly perverse people. The “things” of physics include objects like electrons, and while there are all sorts of complexities that arise in physics, electrons don’t have minds of their own. They don’t have opinions, they don’t differ from each other in weird and arbitrary ways, they don’t get bored in the middle of an experiment, and they don’t get angry at the experimenter and then deliberately try to sabotage

the data set (not that I've ever done that . . .). At a fundamental level psychology is harder than physics.³

Basically, we teach statistics to you as psychologists because you need to be better at stats than physicists. There's actually a saying used sometimes in physics, to the effect that "if your experiment needs statistics, you should have done a better experiment". They have the luxury of being able to say that because their objects of study are pathetically simple in comparison to the vast mess that confronts social scientists. It's not just psychology, really: most social sciences are desperately reliant on statistics. Not because we're bad experimenters, but because we've picked a harder problem to solve. We teach you stats because you really, really need it.

- **Can't someone else do the statistics?**

To some extent, but not completely. It's true that you don't need to become a fully trained statistician just to do psychology, but you do need to reach a certain level of statistical competence. In my view, there's three reasons that every psychological researcher ought to be able to do basic statistics:

- Firstly, there's the fundamental reason: statistics is deeply intertwined with research design. If you want to be good at designing psychological studies, you need to at least understand the basics of stats.
- Secondly, if you want to be good at the psychological side of the research, then you need to be able to understand the psychological literature, right? But almost every paper in the psychological literature reports the results of statistical analyses. So if you really want to understand the psychology, you need to be able to understand what other people did with their data. And that means understanding a certain amount of statistics.
- Thirdly, there's a big practical problem with being dependent on other people to do all your statistics: statistical analysis is *expensive*. If you ever get bored and want to look up how much the Australian government charges for university fees, you'll notice something interesting: statistics is designated as a "national priority" category, and so the fees are much, much lower than for any other area of study. This is because there's a massive shortage of statisticians out there. So, from your perspective as a psychological researcher, the laws of supply and demand aren't exactly on your side here! As a result, in almost any real life situation where you want to do psychological research, the cruel facts will be that you don't have enough money to afford a statistician. So the economics of the situation mean that you have to be pretty self-sufficient.

Note that a lot of these reasons generalise beyond researchers. If you want to be a practicing psychologist and stay on top of the field, it helps to be able to read the scientific literature, which relies pretty heavily on statistics.

- **I don't care about jobs, research, or clinical work. Do I need statistics?**

Okay, now you're just messing with me. Still, I think it should matter to you too. Statistics should matter to you in the same way that statistics should matter to *everyone*: we live in the 21st century, and data are *everywhere*. Frankly, given the world in which we live these days, a basic knowledge of statistics is pretty damn close to a survival tool! Which is the topic of the next section...

³Which might explain why physics is just a teensy bit further advanced as a science than we are.

1.4

Statistics in everyday life

*“We are drowning in information,
but we are starved for knowledge”*

– Various authors, original probably John Naisbitt

When I started writing up my lecture notes I took the 20 most recent news articles posted to the ABC news website. Of those 20 articles, it turned out that 8 of them involved a discussion of something that I would call a statistical topic; 6 of those made a mistake. The most common error, if you’re curious, was failing to report baseline data (e.g., the article mentions that 5% of people in situation X have some characteristic Y, but doesn’t say how common the characteristic is for everyone else!) The point I’m trying to make here isn’t that journalists are bad at statistics (though they almost always are), it’s that a basic knowledge of statistics is very helpful for trying to figure out when someone else is either making a mistake or even lying to you. In fact, one of the biggest things that a knowledge of statistics does to you is cause you to get angry at the newspaper or the internet on a far more frequent basis: you can find a good example of this in Section 5.1.5. In later versions of this book I’ll try to include more anecdotes along those lines.

1.5

There's more to research methods than statistics

So far, most of what I’ve talked about is statistics, and so you’d be forgiven for thinking that statistics is all I care about in life. To be fair, you wouldn’t be far wrong, but research methodology is a broader concept than statistics. So most research methods courses will cover a lot of topics that relate much more to the pragmatics of research design, and in particular the issues that you encounter when trying to do research with humans. However, about 99% of student *fears* relate to the statistics part of the course, so I’ve focused on the stats in this discussion, and hopefully I’ve convinced you that statistics matters, and more importantly, that it’s not to be feared. That being said, it’s pretty typical for introductory research methods classes to be very stats-heavy. This is not (usually) because the lecturers are evil people. Quite the contrary, in fact. Introductory classes focus a lot on the statistics because you almost always find yourself needing statistics before you need the other research methods training. Why? Because almost all of your assignments in other classes will rely on statistical training, to a much greater extent than they rely on other methodological tools. It’s not common for undergraduate assignments to require you to design your own study from the ground up (in which case you would need to know a lot about research design), but it *is* common for assignments to ask you to analyse and interpret data that were collected in a study that someone else designed (in which case you need statistics). In that sense, from the perspective of allowing you to do well in all your other classes, the statistics is more urgent.

But note that “urgent” is different from “important” – they both matter. I really do want to stress that research design is just as important as data analysis, and this book does spend a fair amount of time on it. However, while statistics has a kind of universality, and provides a set of core tools that are useful for most types of psychological research, the research methods side isn’t quite so universal. There are some general principles that everyone should think about, but a lot of research design is very idiosyncratic, and is specific to the area of research that you want to engage in. To the extent that it’s the details that matter, those details don’t usually show up in an introductory stats and research methods class.

2. A brief introduction to research design

In this chapter, we're going to start thinking about the basic ideas that go into designing a study, collecting data, checking whether your data collection works, and so on. It won't give you enough information to allow you to design studies of your own, but it will give you a lot of the basic tools that you need to assess the studies done by other people. However, since the focus of this book is much more on data analysis than on data collection, I'm only giving a very brief overview. Note that this chapter is "special" in two ways. Firstly, it's much more psychology-specific than the later chapters. Secondly, it focuses much more heavily on the scientific problem of research methodology, and much less on the statistical problem of data analysis. Nevertheless, the two problems are related to one another, so it's traditional for stats textbooks to discuss the problem in a little detail. This chapter relies heavily on [Campbell and Stanley \(1963\)](#) for the discussion of study design, and [Stevens \(1946\)](#) for the discussion of scales of measurement. Later versions will attempt to be more precise in the citations.

2.1

Introduction to psychological measurement

The first thing to understand is data collection can be thought of as a kind of **measurement**. That is, what we're trying to do here is measure something about human behaviour or the human mind. What do I mean by "measurement"?

2.1.1 Some thoughts about psychological measurement

Measurement itself is a subtle concept, but basically it comes down to finding some way of assigning numbers, or labels, or some other kind of well-defined descriptions to "stuff". So, any of the following would count as a psychological measurement:

- My **age** is *33 years*.
- I *do not* like anchovies.
- My **chromosomal gender** is *male*.
- My **self-identified gender** is *male*.

In the short list above, the **bolded part** is "the thing to be measured", and the *italicised part* is "the measurement itself". In fact, we can expand on this a little bit, by thinking about the set of possible measurements that could have arisen in each case:

- My **age** (in years) could have been *0, 1, 2, 3 …*, etc. The upper bound on what my age could possibly be is a bit fuzzy, but in practice you'd be safe in saying that the largest possible age is *150*, since no human has ever lived that long.
- When asked if I like **anchovies**, I might have said that *I do*, or *I do not*, or *I have no opinion*, or *I sometimes do*.
- My **chromosomal gender** is almost certainly going to be *male (XY)* or *female (XX)*, but there are a few other possibilities. I could also have *Klinefelter's syndrome (XXY)*, which is more similar to male than to female. And I imagine there are other possibilities too.
- My **self-identified gender** is also very likely to be *male* or *female*, but it doesn't have to agree with my chromosomal gender. I may also choose to identify with *neither*, or to explicitly call myself *transgender*.

As you can see, for some things (like age) it seems fairly obvious what the set of possible measurements should be, whereas for other things it gets a bit tricky. But I want to point out that even in the case of someone's age, it's much more subtle than this. For instance, in the example above, I assumed that it was okay to measure age in years. But if you're a developmental psychologist, that's way too crude, and so you often measure age in *years and months* (if a child is 2 years and 11 months, this is usually written as "2;11"). If you're interested in newborns, you might want to measure age in *days since birth*, maybe even *hours since birth*. In other words, the way in which you specify the allowable measurement values is important.

Looking at this a bit more closely, you might also realise that the concept of "age" isn't actually all that precise. In general, when we say "age" we implicitly mean "the length of time since birth". But that's not always the right way to do it. Suppose you're interested in how newborn babies control their eye movements. If you're interested in kids that young, you might also start to worry that "birth" is not the only meaningful point in time to care about. If Baby Alice is born 3 weeks premature and Baby Bianca is born 1 week late, would it really make sense to say that they are the "same age" if we encountered them "2 hours after birth"? In one sense, yes: by social convention, we use birth as our reference point for talking about age in everyday life, since it defines the amount of time the person has been operating as an independent entity in the world, but from a scientific perspective that's not the only thing we care about. When we think about the biology of human beings, it's often useful to think of ourselves as organisms that have been growing and maturing since conception, and from that perspective Alice and Bianca aren't the same age at all. So you might want to define the concept of "age" in two different ways: the length of time since conception, and the length of time since birth. When dealing with adults, it won't make much difference, but when dealing with newborns it might.

Moving beyond these issues, there's the question of methodology. What specific "measurement method" are you going to use to find out someone's age? As before, there are lots of different possibilities:

- You could just ask people "how old are you?" The method of self-report is fast, cheap and easy, but it only works with people old enough to understand the question, and some people lie about their age.
- You could ask an authority (e.g., a parent) "how old is your child?" This method is fast, and when dealing with kids it's not all that hard since the parent is almost always around. It doesn't work as well if you want to know "age since conception", since a lot of parents can't say for sure when conception took place. For that, you might need a different authority (e.g., an obstetrician).
- You could look up official records, like birth certificates. This is time consuming and annoying, but it has its uses (e.g., if the person is now dead).

2.1.2 Operationalisation: defining your measurement

All of the ideas discussed in the previous section all relate to the concept of **operationalisation**. To be a bit more precise about the idea, operationalisation is the process by which we take a meaningful but somewhat vague concept, and turn it into a precise measurement. The process of operationalisation can involve several different things:

- Being precise about what you are trying to measure. For instance, does “age” mean “time since birth” or “time since conception” in the context of your research?
- Determining what method you will use to measure it. Will you use self-report to measure age, ask a parent, or look up an official record? If you’re using self-report, how will you phrase the question?
- Defining the set of the allowable values that the measurement can take. Note that these values don’t always have to be numerical, though they often are. When measuring age, the values are numerical, but we still need to think carefully about what numbers are allowed. Do we want age in years, years and months, days, hours? Etc. For other types of measurements (e.g., gender), the values aren’t numerical. But, just as before, we need to think about what values are allowed. If we’re asking people to self-report their gender, what options do we allow them to choose between? Is it enough to allow only “male” or “female”? Do you need an “other” option? Or should we not give people any specific options, and let them answer in their own words? And if you open up the set of possible values to include all verbal responses, how will you interpret their answers?

Operationalisation is a tricky business, and there’s no “one, true way” to do it. The way in which you choose to operationalise the informal concept of “age” or “gender” into a formal measurement depends on what you need to use the measurement for. Often you’ll find that the community of scientists who work in your area have some fairly well-established ideas for how to go about it. In other words, operationalisation needs to be thought through on a case by case basis. Nevertheless, while there are a lot of issues that are specific to each individual research project, there are some aspects to it that are pretty general.

Before moving on, I want to take a moment to clear up our terminology, and in the process introduce one more term. Here are four different things that are closely related to each other:

- **A theoretical construct.** This is the thing that you’re trying to take a measurement of, like “age”, “gender” or an “opinion”. A theoretical construct can’t be directly observed, and often they’re actually a bit vague.
- **A measure.** The measure refers to the method or the tool that you use to make your observations. A question in a survey, a behavioural observation or a brain scan could all count as a measure.
- **An operationalisation.** The term “operationalisation” refers to the logical connection between the measure and the theoretical construct, or to the process by which we try to derive a measure from a theoretical construct.
- **A variable.** Finally, a new term. A variable is what we end up with when we apply our measure to something in the world. That is, variables are the actual “data” that we end up with in our data sets.

In practice, even scientists tend to blur the distinction between these things, but it’s very helpful to try to understand the differences.

2.2

Scales of measurement

As the previous section indicates, the outcome of a psychological measurement is called a variable. But not all variables are of the same qualitative type, and it's very useful to understand what types there are. A very useful concept for distinguishing between different types of variables is what's known as **scales of measurement**.

2.2.1 Nominal scale

A **nominal scale** variable is one in which there is no particular relationship between the different possibilities: for these kinds of variables it doesn't make any sense to say that one of them is "bigger" or "better" than any other one, and it absolutely doesn't make any sense to average them. The classic example for this is "eye colour". Eyes can be blue, green and brown, among other possibilities, but none of them is any "better" than any other one. As a result, it would feel really weird to talk about an "average eye colour". Similarly, gender is nominal too: male isn't better or worse than female, neither does it make sense to try to talk about an "average gender". In short, nominal scale variables are those for which the only thing you can say about the different possibilities is that they are different. That's it.

Let's take a slightly closer look at this. Suppose I was doing research on how people commute to and from work. One variable I would have to measure would be what kind of transportation people use to get to work. This "transport type" variable could have quite a few possible values, including: "train", "bus", "car", "bicycle", etc. For now, let's suppose that these four are the only possibilities, and suppose that when I ask 100 people how they got to work today, and I get this:

Transportation	Number of people
(1) Train	12
(2) Bus	30
(3) Car	48
(4) Bicycle	10

So, what's the average transportation type? Obviously, the answer here is that there isn't one. It's a silly question to ask. You can say that travel by car is the most popular method, and travel by train is the least popular method, but that's about all. Similarly, notice that the order in which I list the options isn't very interesting. I could have chosen to display the data like this

Transportation	Number of people
(3) Car	48
(1) Train	12
(4) Bicycle	10
(2) Bus	30

and nothing really changes.

2.2.2 Ordinal scale

Ordinal scale variables have a bit more structure than nominal scale variables, but not by a lot. An ordinal scale variable is one in which there is a natural, meaningful way to order the different possibilities,

but you can't do anything else. The usual example given of an ordinal variable is "finishing position in a race". You *can* say that the person who finished first was faster than the person who finished second, but you *don't* know how much faster. As a consequence we know that 1st > 2nd, and we know that 2nd > 3rd, but the difference between 1st and 2nd might be much larger than the difference between 2nd and 3rd.

Here's an more psychologically interesting example. Suppose I'm interested in people's attitudes to climate change, and I ask them to pick one of these four statements that most closely matches their beliefs:

- (1) Temperatures are rising, because of human activity
- (2) Temperatures are rising, but we don't know why
- (3) Temperatures are rising, but not because of humans
- (4) Temperatures are not rising

Notice that these four statements actually do have a natural ordering, in terms of "the extent to which they agree with the current science". Statement 1 is a close match, statement 2 is a reasonable match, statement 3 isn't a very good match, and statement 4 is in strong opposition to the science. So, in terms of the thing I'm interested in (the extent to which people endorse the science), I can order the items as 1 > 2 > 3 > 4. Since this ordering exists, it would be very weird to list the options like this...

- (3) Temperatures are rising, but not because of humans
- (1) Temperatures are rising, because of human activity
- (4) Temperatures are not rising
- (2) Temperatures are rising, but we don't know why

...because it seems to violate the natural "structure" to the question.

So, let's suppose I asked 100 people these questions, and got the following answers:

Response	Number
(1) Temperatures are rising, because of human activity	51
(2) Temperatures are rising, but we don't know why	20
(3) Temperatures are rising, but not because of humans	10
(4) Temperatures are not rising	19

When analysing these data, it seems quite reasonable to try to group (1), (2) and (3) together, and say that 81 of 100 people were willing to *at least partially* endorse the science. And it's *also* quite reasonable to group (2), (3) and (4) together and say that 49 of 100 people registered *at least some disagreement* with the dominant scientific view. However, it would be entirely bizarre to try to group (1), (2) and (4) together and say that 90 of 100 people said... what? There's nothing sensible that allows you to group those responses together at all.

That said, notice that while we *can* use the natural ordering of these items to construct sensible groupings, what we *can't* do is average them. For instance, in my simple example here, the "average" response to the question is 1.97. If you can tell me what that means, I'd love to know. Because that sounds like gibberish to me!

2.2.3 Interval scale

In contrast to nominal and ordinal scale variables, **interval scale** and ratio scale variables are variables for which the numerical value is genuinely meaningful. In the case of interval scale variables, the *differences* between the numbers are interpretable, but the variable doesn't have a "natural" zero value.

A good example of an interval scale variable is measuring temperature in degrees celsius. For instance, if it was 15° yesterday and 18° today, then the 3° difference between the two is genuinely meaningful. Moreover, that 3° difference is *exactly the same* as the 3° difference between 7° and 10° . In short, addition and subtraction are meaningful for interval scale variables.

However, notice that the 0° does not mean “no temperature at all”: it actually means “the temperature at which water freezes”, which is pretty arbitrary. As a consequence, it becomes pointless to try to multiply and divide temperatures. It is wrong to say that 20° is *twice as hot* as 10° , just as it is weird and meaningless to try to claim that 20° is negative two times as hot as -10° .

Again, lets look at a more psychological example. Suppose I’m interested in looking at how the attitudes of first-year university students have changed over time. Obviously, I’m going to want to record the year in which each student started. This is an interval scale variable. A student who started in 2003 did arrive 5 years before a student who started in 2008. However, it would be completely insane for me to divide 2008 by 2003 and say that the second student started “1.0024 times later” than the first one. That doesn’t make any sense at all.

2.2.4 Ratio scale

The fourth and final type of variable to consider is a **ratio scale** variable, in which zero really means zero, and it’s okay to multiply and divide. A good psychological example of a ratio scale variable is response time (RT). In a lot of tasks it’s very common to record the amount of time somebody takes to solve a problem or answer a question, because it’s an indicator of how difficult the task is. Suppose that Alan takes 2.3 seconds to respond to a question, whereas Ben takes 3.1 seconds. As with an interval scale variable, addition and subtraction are both meaningful here. Ben really did take $3.1 - 2.3 = 0.8$ seconds longer than Alan did. However, notice that multiplication and division also make sense here too: Ben took $3.1/2.3 = 1.35$ times as long as Alan did to answer the question. And the reason why you can do this is that, for a ratio scale variable such as RT, “zero seconds” really does mean “no time at all”.

2.2.5 Continuous versus discrete variables

There’s a second kind of distinction that you need to be aware of, regarding what types of variables you can run into. This is the distinction between continuous variables and discrete variables. The difference between these is as follows:

- A **continuous variable** is one in which, for any two values that you can think of, it’s always logically possible to have another value in between.
- A **discrete variable** is, in effect, a variable that isn’t continuous. For a discrete variable, it’s sometimes the case that there’s nothing in the middle.

These definitions probably seem a bit abstract, but they’re pretty simple once you see some examples. For instance, response time is continuous. If Alan takes 3.1 seconds and Ben takes 2.3 seconds to respond to a question, then it’s possible for Cameron’s response time to lie in between, by taking 3.0 seconds. And of course it would also be possible for David to take 3.031 seconds to respond, meaning that his RT would lie in between Cameron’s and Alan’s. And while in practice it might be impossible to measure RT that precisely, it’s certainly possible in principle. Because we can always find a new value for RT in between any two other ones, we say that RT is continuous.

Discrete variables occur when this rule is violated. For example, nominal scale variables are always discrete: there isn’t a type of transportation that falls “in between” trains and bicycles, not in the strict mathematical way that 2.3 falls in between 2 and 3. So transportation type is discrete. Similarly, ordinal scale variables are always discrete: although “2nd place” does fall between “1st place” and “3rd place”, there’s nothing that can logically fall in between “1st place” and “2nd place”. Interval scale and ratio

Table 2.1: The relationship between the scales of measurement and the discrete/continuity distinction. Cells with a tick mark correspond to things that are possible.

	continuous	discrete
nominal		✓
ordinal		✓
interval	✓	✓
ratio	✓	✓

.....

scale variables can go either way. As we saw above, response time (a ratio scale variable) is continuous. Temperature in degrees celsius (an interval scale variable) is also continuous. However, the year you went to school (an interval scale variable) is discrete. There's no year in between 2002 and 2003. The number of questions you get right on a true-or-false test (a ratio scale variable) is also discrete: since a true-or-false question doesn't allow you to be "partially correct", there's nothing in between 5/10 and 6/10. Table 2.1 summarises the relationship between the scales of measurement and the discrete/continuity distinction. Cells with a tick mark correspond to things that are possible. I'm trying to hammer this point home, because (a) some textbooks get this wrong, and (b) people very often say things like "discrete variable" when they mean "nominal scale variable". It's very unfortunate.

2.2.6 Some complexities

Okay, I know you're going to be shocked to hear this, but ... the real world is much messier than this little classification scheme suggests. Very few variables in real life actually fall into these nice neat categories, so you need to be kind of careful not to treat the scales of measurement as if they were hard and fast rules. It doesn't work like that: they're guidelines, intended to help you think about the situations in which you should treat different variables differently. Nothing more.

So let's take a classic example, maybe *the* classic example, of a psychological measurement tool: the **Likert scale**. The humble Likert scale is the bread and butter tool of all survey design. You yourself have filled out hundreds, maybe thousands of them, and odds are you've even used one yourself. Suppose we have a survey question that looks like this:

Which of the following best describes your opinion of the statement that "all pirates are freaking awesome" ...

and then the options presented to the participant are these:

- (1) Strongly disagree
- (2) Disagree
- (3) Neither agree nor disagree
- (4) Agree
- (5) Strongly agree

This set of items is an example of a 5-point Likert scale: people are asked to choose among one of several (in this case 5) clearly ordered possibilities, generally with a verbal descriptor given in each case. However, it's not necessary that all items be explicitly described. This is a perfectly good example of a 5-point Likert scale too:

- (1) Strongly disagree
- (2)

- (3)
- (4)
- (5) Strongly agree

Likert scales are very handy, if somewhat limited, tools. The question is, what kind of variable are they? They're obviously discrete, since you can't give a response of 2.5. They're obviously not nominal scale, since the items are ordered; and they're not ratio scale either, since there's no natural zero.

But are they ordinal scale or interval scale? One argument says that we can't really prove that the difference between "strongly agree" and "agree" is of the same size as the difference between "agree" and "neither agree nor disagree". In fact, in everyday life it's pretty obvious that they're not the same at all. So this suggests that we ought to treat Likert scales as ordinal variables. On the other hand, in practice most participants do seem to take the whole "on a scale from 1 to 5" part fairly seriously, and they tend to act as if the differences between the five response options were fairly similar to one another. As a consequence, a lot of researchers treat Likert scale data as if it were interval scale. It's not interval scale, but in practice it's close enough that we usually think of it as being **quasi-interval scale**.

2.3

Assessing the reliability of a measurement

At this point we've thought a little bit about how to operationalise a theoretical construct and thereby create a psychological measure; and we've seen that by applying psychological measures we end up with variables, which can come in many different types. At this point, we should start discussing the obvious question: is the measurement any good? We'll do this in terms of two related ideas: *reliability* and *validity*. Put simply, the **reliability** of a measure tells you how *precisely* you are measuring something, whereas the validity of a measure tells you how *accurate* the measure is. In this section I'll talk about reliability; we'll talk about validity in the next chapter.

Reliability is actually a very simple concept: it refers to the repeatability or consistency of your measurement. The measurement of my weight by means of a "bathroom scale" is very reliable: if I step on and off the scales over and over again, it'll keep giving me the same answer. Measuring my intelligence by means of "asking my mum" is very unreliable: some days she tells me I'm a bit thick, and other days she tells me I'm a complete moron. Notice that this concept of reliability is different to the question of whether the measurements are correct (the correctness of a measurement relates to its validity). If I'm holding a sack of potatoes when I step on and off of the bathroom scales, the measurement will still be reliable: it will always give me the same answer. However, this highly reliable answer doesn't match up to my true weight at all, therefore it's wrong. In technical terms, this is a *reliable but invalid* measurement. Similarly, while my mum's estimate of my intelligence is a bit unreliable, she might be right. Maybe I'm just not too bright, and so while her estimate of my intelligence fluctuates pretty wildly from day to day, it's basically right. So that would be an *unreliable but valid* measure. Of course, to some extent, notice that if my mum's estimates are too unreliable, it's going to be very hard to figure out which one of her many claims about my intelligence is actually the right one. To some extent, then, a very unreliable measure tends to end up being invalid for practical purposes; so much so that many people would say that reliability is necessary (but not sufficient) to ensure validity.

Okay, now that we're clear on the distinction between reliability and validity, let's have a think about the different ways in which we might measure reliability:

- **Test-retest reliability.** This relates to consistency over time: if we repeat the measurement at a later date, do we get the same answer?

- **Inter-rater reliability.** This relates to consistency across people: if someone else repeats the measurement (e.g., someone else rates my intelligence) will they produce the same answer?
- **Parallel forms reliability.** This relates to consistency across theoretically-equivalent measurements: if I use a different set of bathroom scales to measure my weight, does it give the same answer?
- **Internal consistency reliability.** If a measurement is constructed from lots of different parts that perform similar functions (e.g., a personality questionnaire result is added up across several questions) do the individual parts tend to give similar answers?

Not all measurements need to possess all forms of reliability. For instance, educational assessment can be thought of as a form of measurement. One of the subjects that I teach, *Computational Cognitive Science*, has an assessment structure that has a research component and an exam component (plus other things). The exam component is *intended* to measure something different from the research component, so the assessment as a whole has low internal consistency. However, within the exam there are several questions that are intended to (approximately) measure the same things, and those tend to produce similar outcomes; so the exam on its own has a fairly high internal consistency. Which is as it should be. You should only demand reliability in those situations where you want to be measure the same thing!

2.4

The “role” of variables: predictors and outcomes

Okay, I've got one last piece of terminology that I need to explain to you before moving away from variables. Normally, when we do some research we end up with lots of different variables. Then, when we analyse our data we usually try to explain some of the variables in terms of some of the other variables. It's important to keep the two roles “thing doing the explaining” and “thing being explained” distinct. So let's be clear about this now. Firstly, we might as well get used to the idea of using mathematical symbols to describe variables, since it's going to happen over and over again. Let's denote the “to be explained” variable Y , and denote the variables “doing the explaining” as X_1 , X_2 , etc.

Now, when we doing an analysis, we have different names for X and Y , since they play different roles in the analysis. The classical names for these roles are **independent variable** (IV) and **dependent variable** (DV). The IV is the variable that you use to do the explaining (i.e., X) and the DV is the variable being explained (i.e., Y). The logic behind these names goes like this: if there really is a relationship between X and Y then we can say that Y depends on X , and if we have designed our study “properly” then X isn't dependent on anything else. However, I personally find those names horrible: they're hard to remember and they're highly misleading, because (a) the IV is never actually “independent of everything else” and (b) if there's no relationship, then the DV doesn't actually depend on the IV. And in fact, because I'm not the only person who thinks that IV and DV are just awful names, there are a number of alternatives that I find more appealing. The terms that I'll use in these notes are **predictors** and **outcomes**. The idea here is that what you're trying to do is use X (the predictors) to make guesses about Y (the outcomes).¹ This is summarised in Table 2.2.

¹ Annoyingly, though, there's a lot of different names used out there. I won't list all of them – there would be no point in doing that – other than to note that R often uses “response variable” where I've used “outcome”, and a traditionalist would use “dependent variable”. Sigh. This sort of terminological confusion is very common, I'm afraid.

Table 2.2: The terminology used to distinguish between different roles that a variable can play when analysing a data set. Note that this book will tend to avoid the classical terminology in favour of the newer names.

role of the variable	classical name	modern name
“to be explained”	dependent variable (DV)	outcome
“to do the explaining”	independent variable (IV)	predictor

2.5

Experimental and non-experimental research

One of the big distinctions that you should be aware of is the distinction between “experimental research” and “non-experimental research”. When we make this distinction, what we’re really talking about is the degree of control that the researcher exercises over the people and events in the study.

2.5.1 Experimental research

The key features of **experimental research** is that the researcher controls all aspects of the study, especially what participants experience during the study. In particular, the researcher manipulates or varies the predictor variables (IVs), and then allows the outcome variable (DV) to vary naturally. The idea here is to deliberately vary the predictors (IVs) to see if they have any causal effects on the outcomes. Moreover, in order to ensure that there’s no chance that something other than the predictor variables is causing the outcomes, everything else is kept constant or is in some other way “balanced” to ensure that they have no effect on the results. In practice, it’s almost impossible to *think* of everything else that might have an influence on the outcome of an experiment, much less keep it constant. The standard solution to this is **randomisation**: that is, we randomly assign people to different groups, and then give each group a different treatment (i.e., assign them different values of the predictor variables). We’ll talk more about randomisation later in this course, but for now, it’s enough to say that what randomisation does is minimise (but not eliminate) the chances that there are any systematic difference between groups.

Let’s consider a very simple, completely unrealistic and grossly unethical example. Suppose you wanted to find out if smoking causes lung cancer. One way to do this would be to find people who smoke and people who don’t smoke, and look to see if smokers have a higher rate of lung cancer. This is *not* a proper experiment, since the researcher doesn’t have a lot of control over who is and isn’t a smoker. And this really matters: for instance, it might be that people who choose to smoke cigarettes also tend to have poor diets, or maybe they tend to work in asbestos mines, or whatever. The point here is that the groups (smokers and non-smokers) actually differ on lots of things, not *just* smoking. So it might be that the higher incidence of lung cancer among smokers is caused by something else, not by smoking per se. In technical terms, these other things (e.g. diet) are called “confounds”, and we’ll talk about those in just a moment.

In the meantime, let’s now consider what a proper experiment might look like. Recall that our concern was that smokers and non-smokers might differ in lots of ways. The solution, as long as you have no ethics, is to *control* who smokes and who doesn’t. Specifically, if we randomly divide participants into two groups, and force half of them to become smokers, then it’s very unlikely that the groups will differ in any respect other than the fact that half of them smoke. That way, if our smoking group gets cancer

at a higher rate than the non-smoking group, then we can feel pretty confident that (a) smoking does cause cancer and (b) we're murderers.

2.5.2 Non-experimental research

Non-experimental research is a broad term that covers “any study in which the researcher doesn’t have quite as much control as they do in an experiment”. Obviously, control is something that scientists like to have, but as the previous example illustrates, there are lots of situations in which you can’t or shouldn’t try to obtain that control. Since it’s grossly unethical (and almost certainly criminal) to force people to smoke in order to find out if they get cancer, this is a good example of a situation in which you really shouldn’t try to obtain experimental control. But there are other reasons too. Even leaving aside the ethical issues, our “smoking experiment” does have a few other issues. For instance, when I suggested that we “force” half of the people to become smokers, I must have been talking about *starting* with a sample of non-smokers, and then forcing them to become smokers. While this sounds like the kind of solid, evil experimental design that a mad scientist would love, it might not be a very sound way of investigating the effect in the real world. For instance, suppose that smoking only causes lung cancer when people have poor diets, and suppose also that people who normally smoke do tend to have poor diets. However, since the “smokers” in our experiment aren’t “natural” smokers (i.e., we forced non-smokers to become smokers; they didn’t take on all of the other normal, real life characteristics that smokers might tend to possess) they probably have better diets. As such, in this silly example they wouldn’t get lung cancer, and our experiment will fail, because it violates the structure of the “natural” world (the technical name for this is an “artifactual” result; see later).

One distinction worth making between two types of non-experimental research is the difference between **quasi-experimental research** and **case studies**. The example I discussed earlier – in which we wanted to examine incidence of lung cancer among smokers and non-smokers, without trying to control who smokes and who doesn’t – is a quasi-experimental design. That is, it’s the same as an experiment, but we don’t control the predictors (IVs). We can still use statistics to analyse the results, it’s just that we have to be a lot more careful.

The alternative approach, case studies, aims to provide a very detailed description of one or a few instances. In general, you can’t use statistics to analyse the results of case studies, and it’s usually very hard to draw any general conclusions about “people in general” from a few isolated examples. However, case studies are very useful in some situations. Firstly, there are situations where you don’t have any alternative: neuropsychology has this issue a lot. Sometimes, you just can’t find a lot of people with brain damage in a specific area, so the only thing you can do is describe those cases that you do have in as much detail and with as much care as you can. However, there’s also some genuine advantages to case studies: because you don’t have as many people to study, you have the ability to invest lots of time and effort trying to understand the specific factors at play in each case. This is a very valuable thing to do. As a consequence, case studies can complement the more statistically-oriented approaches that you see in experimental and quasi-experimental designs. We won’t talk much about case studies in these lectures, but they are nevertheless very valuable tools!

2.6

Assessing the validity of a study

More than any other thing, a scientist wants their research to be “valid”. The conceptual idea behind **validity** is very simple: can you trust the results of your study? If not, the study is invalid. However, while it’s easy to state, in practice it’s much harder to check validity than it is to check reliability. And in all honesty, there’s no precise, clearly agreed upon notion of what validity actually is. In fact, there’s

lots of different kinds of validity, each of which raises its own issues, and not all forms of validity are relevant to all studies. I'm going to talk about five different types:

- Internal validity
- External validity
- Construct validity
- Face validity
- Ecological validity

To give you a quick guide as to what matters here... (1) Internal and external validity are the most important, since they tie directly to the fundamental question of whether your study really works. (2) Construct validity asks whether you're measuring what you think you are. (3) Face validity isn't terribly important except insofar as you care about "appearances". (4) Ecological validity is a special case of face validity that corresponds to a kind of appearance that you might care about a lot.

2.6.1 Internal validity

Internal validity refers to the extent to which you are able draw the correct conclusions about the causal relationships between variables. It's called "internal" because it refers to the relationships between things "inside" the study. Let's illustrate the concept with a simple example. Suppose you're interested in finding out whether a university education makes you write better. To do so, you get a group of first year students, ask them to write a 1000 word essay, and count the number of spelling and grammatical errors they make. Then you find some third-year students, who obviously have had more of a university education than the first-years, and repeat the exercise. And let's suppose it turns out that the third-year students produce fewer errors. And so you conclude that a university education improves writing skills. Right? Except... the big problem that you have with this experiment is that the third-year students are older, and they've had more experience with writing things. So it's hard to know for sure what the causal relationship is: Do older people write better? Or people who have had more writing experience? Or people who have had more education? Which of the above is the true *cause* of the superior performance of the third-years? Age? Experience? Education? You can't tell. This is an example of a failure of internal validity, because your study doesn't properly tease apart the *causal* relationships between the different variables.

2.6.2 External validity

External validity relates to the **generalisability** of your findings. That is, to what extent do you expect to see the same pattern of results in "real life" as you saw in your study. To put it a bit more precisely, any study that you do in psychology will involve a fairly specific set of questions or tasks, will occur in a specific environment, and will involve participants that are drawn from a particular subgroup. So, if it turns out that the results don't actually generalise to people and situations beyond the ones that you studied, then what you've got is a lack of external validity.

The classic example of this issue is the fact that a very large proportion of studies in psychology will use undergraduate psychology students as the participants. Obviously, however, the researchers don't care *only* about psychology students; they care about people in general. Given that, a study that uses only psych students as participants always carries a risk of lacking external validity. That is, if there's something "special" about psychology students that makes them different to the general populace in some *relevant* respect, then we may start worrying about a lack of external validity.

That said, it is absolutely critical to realise that a study that uses only psychology students does not necessarily have a problem with external validity. I'll talk about this again later, but it's such a common mistake that I'm going to mention it here. The external validity is threatened by the choice of population if (a) the population from which you sample your participants is very narrow (e.g., psych students), and (b) the narrow population that you sampled from is systematically different from the general population, *in some respect that is relevant to the psychological phenomenon that you intend to study*. The italicised part is the bit that lots of people forget: it is true that psychology undergraduates differ from the general population in lots of ways, and so a study that uses only psych students *may* have problems with external validity. However, if those differences aren't very relevant to the phenomenon that you're studying, then there's nothing to worry about. To make this a bit more concrete, here's two extreme examples:

- You want to measure "attitudes of the general public towards psychotherapy", but all of your participants are psychology students. This study would almost certainly have a problem with external validity.
- You want to measure the effectiveness of a visual illusion, and your participants are all psychology students. This study is very unlikely to have a problem with external validity

Having just spent the last couple of paragraphs focusing on the choice of participants (since that's the big issue that everyone tends to worry most about), it's worth remembering that external validity is a broader concept. The following are also examples of things that might pose a threat to external validity, depending on what kind of study you're doing:

- People might answer a "psychology questionnaire" in a manner that doesn't reflect what they would do in real life.
- Your lab experiment on (say) "human learning" has a different structure to the learning problems people face in real life.

2.6.3 Construct validity

Construct validity is basically a question of whether you're measuring what you want to be measuring. A measurement has good construct validity if it is actually measuring the correct theoretical construct, and bad construct validity if it doesn't. To give very simple (if ridiculous) example, suppose I'm trying to investigate the rates with which university students cheat on their exams. And the way I attempt to measure it is by asking the cheating students to stand up in the lecture theatre so that I can count them. When I do this with a class of 300 students, 0 people claim to be cheaters. So I therefore conclude that the proportion of cheaters in my class is 0%. Clearly this is a bit ridiculous. But the point here is not that this is a very deep methodological example, but rather to explain what construct validity is. The problem with my measure is that while I'm *trying* to measure "the proportion of people who cheat" what I'm actually measuring is "the proportion of people stupid enough to own up to cheating, or bloody minded enough to pretend that they do". Obviously, these aren't the same thing! So my study has gone wrong, because my measurement has very poor construct validity.

2.6.4 Face validity

Face validity simply refers to whether or not a measure "looks like" it's doing what it's supposed to, nothing more. If I design a test of intelligence, and people look at it and they say "no, that test doesn't measure intelligence", then the measure lacks face validity. It's as simple as that. Obviously, face validity isn't very important from a pure scientific perspective. After all, what we care about is whether or not

the measure *actually* does what it's supposed to do, not whether it *looks like* it does what it's supposed to do. As a consequence, we generally don't care very much about face validity. That said, the concept of face validity serves three useful pragmatic purposes:

- Sometimes, an experienced scientist will have a “hunch” that a particular measure won’t work. While these sorts of hunches have no strict evidentiary value, it’s often worth paying attention to them. Because often times people have knowledge that they can’t quite verbalise, so there might be something to worry about even if you can’t quite say why. In other words, when someone you trust criticises the face validity of your study, it’s worth taking the time to think more carefully about your design to see if you can think of reasons why it might go awry. Mind you, if you don’t find any reason for concern, then you should probably not worry: after all, face validity really doesn’t matter much.
- Often (very often), completely uninformed people will also have a “hunch” that your research is crap. And they’ll criticise it on the internet or something. On close inspection, you’ll often notice that these criticisms are actually focused entirely on how the study “looks”, but not on anything deeper. The concept of face validity is useful for gently explaining to people that they need to substantiate their arguments further.
- Expanding on the last point, if the beliefs of untrained people are critical (e.g., this is often the case for applied research where you actually want to convince policy makers of something or other) then you *have* to care about face validity. Simply because – whether you like it or not – a lot of people will use face validity as a proxy for real validity. If you want the government to change a law on scientific, psychological grounds, then it won’t matter how good your studies “really” are. If they lack face validity, you’ll find that politicians ignore you. Of course, it’s somewhat unfair that policy often depends more on appearance than fact, but that’s how things go.

2.6.5 Ecological validity

Ecological validity is a different notion of validity, which is similar to external validity, but less important. The idea is that, in order to be ecologically valid, the entire set up of the study should closely approximate the real world scenario that is being investigated. In a sense, ecological validity is a kind of face validity – it relates mostly to whether the study “looks” right, but with a bit more rigour to it. To be ecologically valid, the study has to look right in a fairly specific way. The idea behind it is the intuition that a study that is ecologically valid is more likely to be externally valid. It’s no guarantee, of course. But the nice thing about ecological validity is that it’s much easier to check whether a study is ecologically valid than it is to check whether a study is externally valid. An simple example would be eyewitness identification studies. Most of these studies tend to be done in a university setting, often with fairly simple array of faces to look at rather than a line up. The length of time between seeing the “criminal” and being asked to identify the suspect in the “line up” is usually shorter. The “crime” isn’t real, so there’s no chance that the witness being scared, and there’s no police officers present, so there’s not as much chance of feeling pressured. These things all mean that the study *definitely* lacks ecological validity. They might (but might not) mean that it also lacks external validity.

2.7 _____

Confounds, artifacts and other threats to validity

If we look at the issue of validity in the most general fashion, the two biggest worries that we have are *confounds* and *artifact*. These two terms are defined in the following way:

- **Confound:** A confound is an additional, often unmeasured variable² that turns out to be related to both the predictors and the outcomes. The existence of confounds threatens the internal validity of the study because you can't tell whether the predictor causes the outcome, or if the confounding variable causes it, etc.
- **Artifact:** A result is said to be “artifactual” if it only holds in the special situation that you happened to test in your study. The possibility that your result is an artifact describes a threat to your external validity, because it raises the possibility that you can't generalise your results to the actual population that you care about.

As a general rule confounds are a bigger concern for non-experimental studies, precisely because they're not proper experiments: by definition, you're leaving lots of things uncontrolled, so there's a lot of scope for confounds working their way into your study. Experimental research tends to be much less vulnerable to confounds: the more control you have over what happens during the study, the more you can prevent confounds from appearing.

However, there's always swings and roundabouts, and when we start thinking about artifacts rather than confounds, the shoe is very firmly on the other foot. For the most part, artifactual results tend to be a concern for experimental studies than for non-experimental studies. To see this, it helps to realise that the reason that a lot of studies are non-experimental is precisely because what the researcher is trying to do is examine human behaviour in a more naturalistic context. By working in a more real-world context, you lose experimental control (making yourself vulnerable to confounds) but because you tend to be studying human psychology “in the wild” you reduce the chances of getting an artifactual result. Or, to put it another way, when you take psychology out of the wild and bring it into the lab (which we usually have to do to gain our experimental control), you always run the risk of accidentally studying something different than you wanted to study: which is more or less the definition of an artifact.

Be warned though: the above is a rough guide only. It's absolutely possible to have confounds in an experiment, and to get artifactual results with non-experimental studies. This can happen for all sorts of reasons, not least of which is researcher error. In practice, it's really hard to think everything through ahead of time, and even very good researchers make mistakes. But other times it's unavoidable, simply because the researcher has ethics (e.g., see “differential attrition”).

Okay. There's a sense in which almost any threat to validity can be characterised as a confound or an artifact: they're pretty vague concepts. So let's have a look at some of the most common examples...

2.7.1 History effects

History effects refer to the possibility that specific events may occur during the study itself that might influence the outcomes. For instance, something might happen in between a pre-test and a post-test. Or, in between testing participant 23 and participant 24. Alternatively, it might be that you're looking at an older study, which was perfectly valid for its time, but the world has changed enough since then that the conclusions are no longer trustworthy. Examples of things that would count as history effects:

- You're interested in how people think about risk and uncertainty. You started your data collection in December 2010. But finding participants and collecting data takes time, so you're still finding new people in February 2011. Unfortunately for you (and even more unfortunately for others), the Queensland floods occurred in January 2011, causing billions of dollars of damage and killing

²The reason why I say that it's unmeasured is that if you *have* measured it, then you can use some fancy statistical tricks to deal with the confound. Because of the existence of these statistical solutions to the problem of confounds, we often refer to a confound that we have measured and dealt with as a *covariate*. Dealing with covariates is a topic for a more advanced course, but I thought I'd mention it in passing, since it's kind of comforting to at least know that this stuff exists.

many people. Not surprisingly, the people tested in February 2011 express quite different beliefs about handling risk than the people tested in December 2010. Which (if any) of these reflects the “true” beliefs of participants? I think the answer is probably both: the Queensland floods genuinely changed the beliefs of the Australian public, though possibly only temporarily. The key thing here is that the “history” of the people tested in February is quite different to people tested in December.

- You’re testing the psychological effects of a new anti-anxiety drug. So what you do is measure anxiety before administering the drug (e.g., by self-report, and taking physiological measures, let’s say), then you administer the drug, and then you take the same measures afterwards. In the middle, however, because your labs are in Los Angeles, there’s an earthquake, which increases the anxiety of the participants.

2.7.2 Maturation effects

As with history effects, **maturational effects** are fundamentally about change over time. However, maturation effects aren’t in response to specific events. Rather, they relate to how people change on their own over time: we get older, we get tired, we get bored, etc. Some examples of maturation effects:

- When doing developmental psychology research, you need to be aware that children grow up quite rapidly. So, suppose that you want to find out whether some educational trick helps with vocabulary size among 3 year olds. One thing that you need to be aware of is that the vocabulary size of children that age is growing at an incredible rate (multiple words per day), all on its own. If you design your study without taking this maturational effect into account, then you won’t be able to tell if your educational trick works.
- When running a very long experiment in the lab (say, something that goes for 3 hours), it’s very likely that people will begin to get bored and tired, and that this maturational effect will cause performance to decline, regardless of anything else going on in the experiment

2.7.3 Repeated testing effects

An important type of history effect is the effect of **repeated testing**. Suppose I want to take two measurements of some psychological construct (e.g., anxiety). One thing I might be worried about is if the first measurement has an effect on the second measurement. In other words, this is a history effect in which the “event” that influences the second measurement is the first measurement itself! This is not at all uncommon. Examples of this include:

- *Learning and practice*: e.g., “intelligence” at time 2 might appear to go up relative to time 1 because participants learned the general rules of how to solve “intelligence-test-style” questions during the first testing session.
- *Familiarity with the testing situation*: e.g., if people are nervous at time 1, this might make performance go down; after sitting through the first testing situation, they might calm down a lot precisely because they’ve seen what the testing looks like.
- *Auxiliary changes caused by testing*: e.g., if a questionnaire assessing mood is boring, then mood at measurement at time 2 is more likely to become “bored”, precisely because of the boring measurement made at time 1.

2.7.4 Selection bias

Selection bias is a pretty broad term. Suppose that you're running an experiment with two groups of participants, where each group gets a different "treatment", and you want to see if the different treatments lead to different outcomes. However, suppose that, despite your best efforts, you've ended up with a gender imbalance across groups (say, group A has 80% females and group B has 50% females). It might sound like this could never happen, but trust me, it can. This is an example of a selection bias, in which the people "selected into" the two groups have different characteristics. If any of those characteristics turns out to be relevant (say, your treatment works better on females than males) then you're in a lot of trouble.

2.7.5 Differential attrition

One quite subtle danger to be aware of is called **differential attrition**, which is a kind of selection bias that is caused by the study itself. Suppose that, for the first time ever in the history of psychology, I manage to find the perfectly balanced and representative sample of people. I start running "Dan's incredibly long and tedious experiment" on my perfect sample, but then, because my study is incredibly long and tedious, lots of people start dropping out. I can't stop this: as we'll discuss later in the chapter on research ethics, participants absolutely have the right to stop doing any experiment, any time, for whatever reason they feel like, and as researchers we are morally (and professionally) obliged to remind people that they do have this right. So, suppose that "Dan's incredibly long and tedious experiment" has a very high drop out rate. What do you suppose the odds are that this drop out is random? Answer: zero. Almost certainly, the people who remain are more conscientious, more tolerant of boredom etc than those that leave. To the extent that (say) conscientiousness is relevant to the psychological phenomenon that I care about, this attrition can decrease the validity of my results.

When thinking about the effects of differential attrition, it is sometimes helpful to distinguish between two different types. The first is **homogeneous attrition**, in which the attrition effect is the same for all groups, treatments or conditions. In the example I gave above, the differential attrition would be homogeneous if (and only if) the easily bored participants are dropping out of all of the conditions in my experiment at about the same rate. In general, the main effect of homogeneous attrition is likely to be that it makes your sample unrepresentative. As such, the biggest worry that you'll have is that the generalisability of the results decreases: in other words, you lose external validity.

The second type of differential attrition is **heterogeneous attrition**, in which the attrition effect is different for different groups. This is a much bigger problem: not only do you have to worry about your external validity, you also have to worry about your internal validity too. To see why this is the case, let's consider a very dumb study in which I want to see if insulting people makes them act in a more obedient way. Why anyone would actually want to study that I don't know, but let's suppose I really, deeply cared about this. So, I design my experiment with two conditions. In the "treatment" condition, the experimenter insults the participant and then gives them a questionnaire designed to measure obedience. In the "control" condition, the experimenter engages in a bit of pointless chitchat and then gives them the questionnaire. Leaving aside the questionable scientific merits and dubious ethics of such a study, let's have a think about what might go wrong here. As a general rule, when someone insults me to my face, I tend to get much less co-operative. So, there's a pretty good chance that a lot more people are going to drop out of the treatment condition than the control condition. And this drop out isn't going to be random. The people most likely to drop out would probably be the people who don't care all that much about the importance of obediently sitting through the experiment. Since the most bloody minded and disobedient people all left the treatment group but not the control group, we've introduced a confound: the people who actually took the questionnaire in the treatment group were *already* more likely to be dutiful and obedient than the people in the control group. In short, in this study insulting people doesn't make them more obedient: it makes the more disobedient people leave the experiment!

The internal validity of this experiment is completely shot.

2.7.6 Non-response bias

Non-response bias is closely related to selection bias, and to differential attrition. The simplest version of the problem goes like this. You mail out a survey to 1000 people, and only 300 of them reply. The 300 people who replied are almost certainly not a random subsample. People who respond to surveys are systematically different to people who don't. This introduces a problem when trying to generalise from those 300 people who replied, to the population at large; since you now have a very non-random sample. The issue of non-response bias is more general than this, though. Among the (say) 300 people that did respond to the survey, you might find that not everyone answers every question. If (say) 80 people chose not to answer one of your questions, does this introduce problems? As always, the answer is maybe. If the question that wasn't answered was on the last page of the questionnaire, and those 80 surveys were returned with the last page missing, there's a good chance that the missing data isn't a big deal: probably the pages just fell off. However, if the question that 80 people didn't answer was the most confrontational or invasive personal question in the questionnaire, then almost certainly you've got a problem. In essence, what you're dealing with here is what's called the problem of **missing data**. If the data that is missing was "lost" randomly, then it's not a big problem. If it's missing systematically, then it can be a big problem.

2.7.7 Regression to the mean

Regression to the mean is a curious variation on selection bias. It refers to any situation where you select data based on an extreme value on some measure. Because the measure has natural variation, it almost certainly means that when you take a subsequent measurement, that later measurement will be less extreme than the first one, purely by chance.

Here's an example. Suppose I'm interested in whether a psychology education has an adverse effect on very smart kids. To do this, I find the 20 psych I students with the best high school grades and look at how well they're doing at university. It turns out that they're doing a lot better than average, but they're not topping the class at university, even though they did top their classes at high school. What's going on? The natural first thought is that this must mean that the psychology classes must be having an adverse effect on those students. However, while that might very well be the explanation, it's more likely that what you're seeing is an example of "regression to the mean". To see how it works, let's take a moment to think about what is required to get the best mark in a class, regardless of whether that class be at high school or at university. When you've got a big class, there are going to be *lots* of very smart people enrolled. To get the best mark you have to be very smart, work very hard, and be a bit lucky. The exam has to ask just the right questions for your idiosyncratic skills, and you have to not make any dumb mistakes (we all do that sometimes) when answering them. And that's the thing: intelligence and hard work are transferrable from one class to the next. Luck isn't. The people who got lucky in high school won't be the same as the people who get lucky at university. That's the very definition of "luck". The consequence of this is that, when you select people at the very extreme values of one measurement (the top 20 students), you're selecting for hard work, skill and luck. But because the luck doesn't transfer to the second measurement (only the skill and work), these people will all be expected to drop a little bit when you measure them a second time (at university). So their scores fall back a little bit, back towards everyone else. This is regression to the mean.

Regression to the mean is surprisingly common. For instance, if two very tall people have kids, their children will tend to be taller than average, but not as tall as the parents. The reverse happens with very short parents: two very short parents will tend to have short children, but nevertheless those kids will tend to be taller than the parents. It can also be extremely subtle. Quite some time ago (sorry, haven't tracked down the references yet) there was some research done that suggested that people learn better

from negative feedback than from positive feedback. However, the way that people tried to show this was to give people positive reinforcement whenever they did good, and negative reinforcement when they did bad. And what you see is that after the positive reinforcement, people tended to do worse; but after the negative reinforcement they tended to do better. But! Notice that there's a selection bias here: when people do very well, you're selecting for "high" values, and so you should *expect* (because of regression to the mean) that performance on the next trial should be worse, regardless of whether reinforcement is given. Similarly, after a bad trial, people will tend to improve all on their own.

2.7.8 Experimenter bias

Experimenter bias can come in multiple forms. The basic idea is that the experimenter, despite the best of intentions, can accidentally end up influencing the results of the experiment by subtly communicating the "right answer" or the "desired behaviour" to the participants. Typically, this occurs because the experimenter has special knowledge that the participant does not – either the right answer to the questions being asked, or knowledge of the expected pattern of performance for the condition that the participant is in, and so on. The classic example of this happening is the case study of "Clever Hans", which dates back to 1907. Clever Hans was a horse that apparently was able to read and count, and perform other human like feats of intelligence. After Clever Hans became famous, psychologists started examining his behaviour more closely. It turned out that – not surprisingly – Hans didn't know how to do maths. Rather, Hans was responding to the human observers around him. Because they did know how to count, and the horse had learned to change its behaviour when people changed theirs.

The general solution to the problem of experimenter bias is to engage in double blind studies, where neither the experimenter nor the participant knows which condition the participant is in, or knows what the desired behaviour is. This provides a very good solution to the problem, but it's important to recognise that it's not quite ideal, and hard to pull off perfectly. For instance, the obvious way that I could try to construct a double blind study is to have one of my Ph.D. students (one who doesn't know anything about the experiment) run the study. That feels like it should be enough. The only person (me) who knows all the details (e.g., correct answers to the questions, assignments of participants to conditions) has no interaction with the participants, and the person who does all the talking to people (the Ph.D. student) doesn't know anything. Except, that last part is very unlikely to be true. In order for the Ph.D. student to run the study effectively, they need to have been briefed by me, the researcher. And, as it happens, the Ph.D. student also knows me, and knows a bit about my general beliefs about people and psychology (e.g., I tend to think humans are much smarter than psychologists give them credit for). As a result of all this, it's almost impossible for the experimenter to avoid knowing a little bit about what expectations I have. And even a little bit of knowledge can have an effect: suppose the experimenter accidentally conveys the fact that the participants are expected to do well in this task. Well, there's a thing called the "Pygmalion effect": if you expect great things of people, they'll rise to the occasion; but if you expect them to fail, they'll do that too. In other words, the expectations become a self-fulfilling prophecy.

2.7.9 Demand effects and reactivity

When talking about experimenter bias, the worry is that the experimenter's knowledge or desires for the experiment are communicated to the participants, and that these effect people's behaviour. However, even if you manage to stop this from happening, it's almost impossible to stop people from knowing that they're part of a psychological study. And the mere fact of knowing that someone is watching/studying you can have a pretty big effect on behaviour. This is generally referred to as **reactivity** or **demand effects**. The basic idea is captured by the Hawthorne effect: people alter their performance because of the attention that the study focuses on them. The effect takes its name from a the "Hawthorne Works" factory outside of Chicago. A study done in the 1920s looking at the effects of lighting on worker

productivity at the factory turned out to be an effect of the fact that the workers knew they were being studied, rather than the lighting.

To get a bit more specific about some of the ways in which the mere fact of being in a study can change how people behave, it helps to think like a social psychologist and look at some of the *roles* that people might adopt during an experiment, but might not adopt if the corresponding events were occurring in the real world:

- The *good participant* tries to be too helpful to the researcher: he or she seeks to figure out the experimenter's hypotheses and confirm them.
- The *negative participant* does the exact opposite of the good participant: he or she seeks to break or destroy the study or the hypothesis in some way.
- The *faithful participant* is unnaturally obedient: he or she seeks to follow instructions perfectly, regardless of what might have happened in a more realistic setting.
- The *apprehensive participant* gets nervous about being tested or studied, so much so that his or her behaviour becomes highly unnatural, or overly socially desirable.

2.7.10 Placebo effects

The **placebo effect** is a specific type of demand effect that we worry a lot about. It refers to the situation where the mere fact of being treated causes an improvement in outcomes. The classic example comes from clinical trials: if you give people a completely chemically inert drug and tell them that it's a cure for a disease, they will tend to get better faster than people who aren't treated at all. In other words, it is people's belief that they are being treated that causes the improved outcomes, not the drug.

2.7.11 Situation, measurement and subpopulation effects

In some respects, these terms are a catch-all term for "all other threats to external validity". They refer to the fact that the choice of subpopulation from which you draw your participants, the location, timing and manner in which you run your study (including who collects the data) and the tools that you use to make your measurements might all be influencing the results. Specifically, the worry is that these things might be influencing the results in such a way that the results won't generalise to a wider array of people, places and measures.

2.7.12 Fraud, deception and self-deception

One final thing that I feel like I should mention. While reading what the textbooks often have to say about assessing the validity of the study, I couldn't help but notice that they seem to make the assumption that the researcher is honest. I find this hilarious. While the vast majority of scientists are honest, in my experience at least, some are not.³ Not only that, as I mentioned earlier, scientists are not immune to belief bias – it's easy for a researcher to end up deceiving themselves into believing the wrong thing, and this can lead them to conduct subtly flawed research, and then hide those flaws when they write it up. So you need to consider not only the (probably unlikely) possibility of outright fraud, but also the (probably quite common) possibility that the research is unintentionally "slanted". Because

³Some people might argue that if you're not honest then you're not a real scientist. Which does have some truth to it I guess, but that's disingenuous (google the "No true Scotsman" fallacy). The fact is that there are lots of people who are employed ostensibly as scientists, and whose work has all of the trappings of science, but who are outright fraudulent. Pretending that they don't exist by saying that they're not scientists is just childish.

I can't find anything on this in the textbook, here's my own attempt to list a few ways in which these issues can arise are:

- **Data fabrication.** Sometimes, people just make up the data. This is occasionally done with “good” intentions. For instance, the researcher believes that the fabricated data do reflect the truth, and may actually reflect “slightly cleaned up” versions of actual data. On other occasions, the fraud is deliberate and malicious. Some high-profile examples where data fabrication has been alleged or shown include Cyril Burt (a psychologist who is thought to have fabricated some of his data), Andrew Wakefield (who has been accused of fabricating his data connecting the MMR vaccine to autism) and Hwang Woo-suk (who falsified a lot of his data on stem cell research).
- **Hoaxes.** Hoaxes share a lot of similarities with data fabrication, but they differ in the intended purpose. A hoax is often a joke, and many of them are intended to be (eventually) discovered. Often, the point of a hoax is to discredit someone or some field. There's quite a few well known scientific hoaxes that have occurred over the years (e.g., Piltdown man) some of were deliberate attempts to discredit particular fields of research (e.g., the Sokal affair).
- **Data misrepresentation.** While fraud gets most of the headlines, it's much more common in my experience to see data being misrepresented. When I say this, I'm not referring to newspapers getting it wrong (which they do, almost always). I'm referring to the fact that often, the data don't actually say what the researchers think they say. My guess is that, almost always, this isn't the result of deliberate dishonesty, it's due to a lack of sophistication in the data analyses. For instance, think back to the example of Simpson's paradox that I discussed in the beginning of these notes. It's very common to see people present “aggregated” data of some kind; and sometimes, when you dig deeper and find the raw data yourself, you find that the aggregated data tell a different story to the disaggregated data. Alternatively, you might find that some aspect of the data is being hidden, because it tells an inconvenient story (e.g., the researcher might choose not to refer to a particular variable). There's a lot of variants on this; many of which are very hard to detect.
- **Study “misdesign”.** Okay, this one is subtle. Basically, the issue here is that a researcher designs a study that has built-in flaws, and those flaws are never reported in the paper. The data that are reported are completely real, and are correctly analysed, but they are produced by a study that is actually quite wrongly put together. The researcher really wants to find a particular effect, and so the study is set up in such a way as to make it “easy” to (artificially) observe that effect. One sneaky way to do this – in case you're feeling like dabbling in a bit of fraud yourself – is to design an experiment in which it's obvious to the participants what they're “supposed” to be doing, and then let reactivity work its magic for you. If you want, you can add all the trappings of double blind experimentation etc. It won't make a difference, since the study materials themselves are subtly telling people what you want them to do. When you write up the results, the fraud won't be obvious to the reader: what's obvious to the participant when they're in the experimental context isn't always obvious to the person reading the paper. Of course, the way I've described this makes it sound like it's always fraud: probably there are cases where this is done deliberately, but in my experience the bigger concern has been with unintentional misdesign. The researcher *believes* ... and so the study just happens to end up with a built in flaw, and that flaw then magically erases itself when the study is written up for publication.
- **Data mining & post hoc hypothesising.** Another way in which the authors of a study can more or less lie about what they found is by engaging in what's referred to as “data mining”. As we'll discuss later in the class, if you keep trying to analyse your data in lots of different ways, you'll eventually find something that “looks” like a real effect but isn't. This is referred to as “data mining”. It used to be quite rare because data analysis used to take weeks, but now that everyone has very powerful statistical software on their computers, it's becoming very common. Data mining per se isn't “wrong”, but the more that you do it, the bigger the risk you're taking. The thing that

is wrong, and I suspect is very common, is *unacknowledged* data mining. That is, the researcher run every possible analysis known to humanity, finds the one that works, and then pretends that this was the only analysis that they ever conducted. Worse yet, they often “invent” a hypothesis after looking at the data, to cover up the data mining. To be clear: it’s not wrong to change your beliefs after looking at the data, and to reanalyse your data using your new “post hoc” hypotheses. What is wrong (and, I suspect, common) is failing to acknowledge that you’ve done so. If you acknowledge that you did it, then other researchers are able to take your behaviour into account. If you don’t, then they can’t. And that makes your behaviour deceptive. Bad!

- **Publication bias & self-censoring.** Finally, a pervasive bias is “non-reporting” of negative results. This is almost impossible to prevent. Journals don’t publish every article that is submitted to them: they prefer to publish articles that find “something”. So, if 20 people run an experiment looking at whether reading Finnegan’s Wake causes insanity in humans, and 19 of them find that it doesn’t, which one do you think is going to get published? Obviously, it’s the one study that did find that Finnegan’s Wake causes insanity⁴. This is an example of a *publication bias*: since no-one ever published the 19 studies that didn’t find an effect, a naive reader would never know that they existed. Worse yet, most researchers “internalise” this bias, and end up *self-censoring* their research. Knowing that negative results aren’t going to be accepted for publication, they never even try to report them. As a friend of mine says “for every experiment that you get published, you also have 10 failures”. And she’s right. The catch is, while some (maybe most) of those studies are failures for boring reasons (e.g. you stuffed something up) others might be genuine “null” results that you ought to acknowledge when you write up the “good” experiment. And telling which is which is often hard to do.

There’s probably a lot more issues like this to think about, but that’ll do to start with. What I really want to point out is the blindingly obvious truth that real world science is conducted by actual humans, and only the most gullible of people automatically assumes that everyone else is honest and impartial. Actual scientists aren’t *that* naive, but for some idiotic reason the world likes to pretend that we’re all morons when it comes to understanding basic human politics.

2.8 _____

Summary

- Introduction to psychological measurement (Section 2.1)
- Scales of measurement and types of variables (Section 2.2)
- Reliability of a measurement (Section 2.3)
- Terminology: predictors and outcomes (Section 2.4)
- Experimental and non-experimental research designs (Section 2.5)
- Validity and its threats (Section 2.6)

⁴Clearly, the real effect is that only insane people would even try to read Finnegan’s Wake.

Part II.

An introduction to R

3. Getting started with R

Robots are nice to work with. They mind their own business, and they never have anything to say.

—Roger Zelazny

In this chapter I'll discuss how to get started in R. I'll briefly talk about how to download and install R, but most of the chapter will be focused on getting you started typing R commands. Our goal in this chapter is not to learn any statistical concepts: we're just trying to learn the basics of how R works and get comfortable interacting with the system. To do this, we'll spend a bit of time using R as a simple calculator, since that's the easiest thing to do with R. In doing so, you'll get a bit of a feel for what it's like to work in R. From there I'll introduce some very basic programming ideas: in particular, I'll talk about the idea of defining *variables* to store information, and a few things that you can do with these variables.

However, before going into any of the specifics, it's worth talking a little about why you might want to use R at all. Given that you're reading this, you've probably got your own reasons. However, if those reasons are “because that's what my stats class uses”, it might be worth explaining a little why your lecturer has chosen to use R for the class. Of course, I don't really know why *other* people choose R, so I'm really talking about why I use it.

- It's sort of obvious, but worth saying anyway: doing your statistics on a computer is faster, easier and more powerful than doing statistics by hand. Computers excel at mindless repetitive tasks, and a lot of statistical calculations are both mindless and repetitive. For most people, the only reason to ever do statistical calculations with pencil and paper is for learning purposes. In my class I do occasionally suggest doing some calculations that way, but the only real value to it is pedagogical. It does help you to get a “feel” for statistics to do some calculations yourself, so it's worth doing it once. But only once!
- Doing statistics in a spreadsheet (e.g., Microsoft Excel) is generally a bad idea in the long run. Although many people are likely feel more familiar with them, spreadsheets are very limited in terms of what analyses they allow you do. If you get into the habit of trying to do your real life data analysis using spreadsheets, then you've dug yourself into a very deep hole.
- Avoiding proprietary software is a very good idea. There are a lot of commercial packages out there that you can buy, some of which I like and some of which I don't. They're usually very glossy in their appearance, and generally very powerful (much more powerful than spreadsheets). However, they're also very expensive: usually, the company sells “student versions” (crippled versions of the real thing) very cheaply; they sell full powered “educational versions” at a price that makes me wince; and they sell commercial licences with a staggeringly high price tag. The business model here is to suck you in during your student days, and then leave you dependent on their tools when you go out into the real world. It's hard to blame them for trying, but personally I'm not in favour of

shelling out thousands of dollars if I can avoid it. And you can avoid it: if you make use of packages like R that are open source and free, you never get trapped having to pay exorbitant licensing fees.

- Something that you might not appreciate now, but will love later on if you do anything involving data analysis, is the fact that R is highly extensible. When you download and install R, you get all the basic “packages”, and those are very powerful on their own. However, because R is so open and so widely used, it’s become something of a standard tool in statistics, and so lots of people write their own packages that extend the system. And these are freely available too. One of the consequences of this, I’ve noticed, is that if you open up an advanced textbook (a recent one, that is) rather than introductory textbooks, is that a *lot* of them use R. In other words, if you learn how to do your basic statistics in R, then you’re a lot closer to being able to use the state of the art methods than you would be if you’d started out with a “simpler” system: so if you want to become a genuine expert in psychological data analysis, learning R is a very good use of your time.
- Related to the previous point: R is a real programming language. As you get better at using R for data analysis, you’re also learning to program. To some people this might seem like a bad thing, but in truth, programming is a core research skill across a lot of the social and behavioural sciences. Think about how many surveys and experiments are done online, or presented on computers. Think about all those online social environments (e.g., Facebook, World of Warcraft) which you might be interested in studying; and maybe collecting data from in an automated fashion. Think about artificial intelligence systems, computer vision and speech recognition. If any of these are things that you think you might want to be involved in – as someone “doing research in psychology”, that is – you’ll need to know a bit of programming. And if you don’t already know how to program, then learning how to do statistics using R is a nice way to start.

Those are the main reasons I use R. It’s not without its flaws: it’s not easy to learn, and it has a few very annoying quirks to it that we’re all pretty much stuck with, but on the whole I think the strengths outweigh the weakness; more so than any other option I’ve encountered so far.

3.1

Installing R

Okay, enough with the sales pitch. Let’s get started. Just as with any piece of software, R needs to be installed on a “computer”, which is a magical box that does cool things and delivers free ponies. Or something along those lines: I may be confusing computers with the iPad marketing campaigns. Anyway, R is freely distributed online, and you can download it from the R homepage, which is:

<http://cran.r-project.org/>

At the top of the page – under the heading “Download and Install R” – you’ll see separate links for Windows users, Mac users, and Linux users. If you follow the relevant link, you’ll see that the online instructions are pretty self-explanatory, but I’ll walk you through the installation anyway. As of this writing, the current version of R is 2.14.0, but they usually issue (free) updates every six months, so you’ll probably have a newer version.¹

¹Although R is updated frequently, it doesn’t usually make much of a difference for the sort of work we’ll do in this book. In fact, during the writing of the book I upgraded twice, and didn’t have to change anything except these sections describing the downloading.

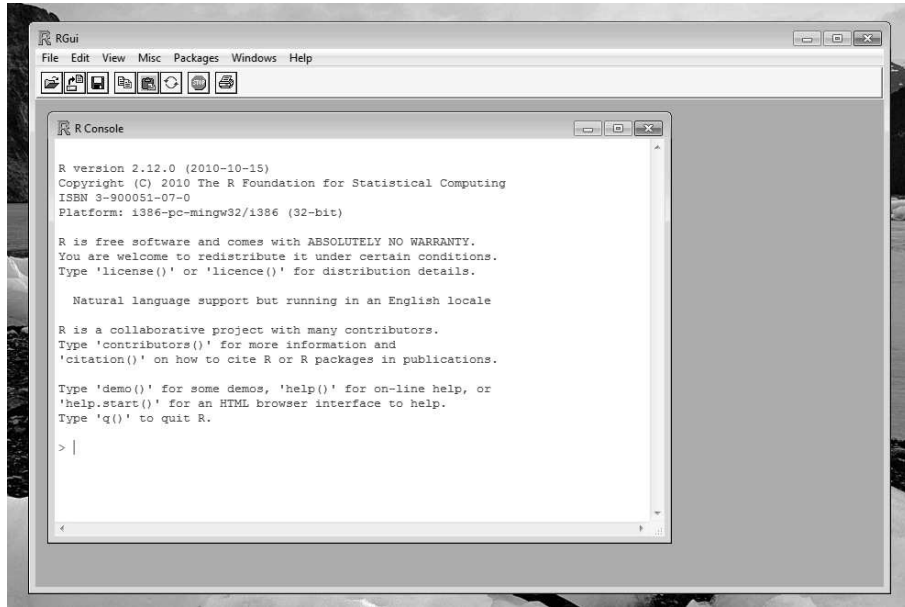


Figure 3.1: A screenshot of R running on a Windows machine. Apparently the version of R I was using when I took this screenshot was 2.12.0, which tells me that I've been working on this book for over a year.

.....

3.1.1 Installing R on a Windows computer

If you're trying to install R on a computer running Windows and follow the links from the CRAN homepage, they should take you to a page that has a link that reads "Download R 2.14.0 for Windows" featuring prominently on the page. When you click on the link, your browser should start downloading a file called `R-2.14.0-win.exe`, or whatever the equivalent version number is by the time you read this. The file for version 2.14.0 is about 45MB in size, so it may take some time depending on how fast your internet connection is. Once you've downloaded the file, double click to install it. As with any software you download online, Windows will ask you some questions about whether you trust the file and so on. After you click through those, it'll ask you where you want to install it, and what components you want to install. The default values should be fine for most people, so again, just click through. Once all that is done, you should have R installed on your system. You can access it from the Start menu, or from the desktop if you asked it to add a shortcut there. When you open up R, you'll see something that looks very similar to the screenshot in Figure 3.1.

3.1.2 Installing R on a Mac

When you click on the Mac OS X link, you should find yourself on a page with the title "R for Mac OS X". The vast majority of Mac users will have a fairly recent version of the operating system: as long as you're running Mac OS X 10.5 (Leopard) or higher, then you'll be fine.² There's a fairly prominent link on the page called "`R-2.14.0.pkg`", which is the one you want if you're running Leopard (10.5), Snow Leopard (10.6) or Lion (10.7). Click on that link and you'll start downloading the installer file, which is

²If you're running an older version of the Mac OS, then you need to follow the link to the "old" page (<http://cran.r-project.org/bin/macosx.old/>). You should be able to find the installer file that you need at the bottom of the page.

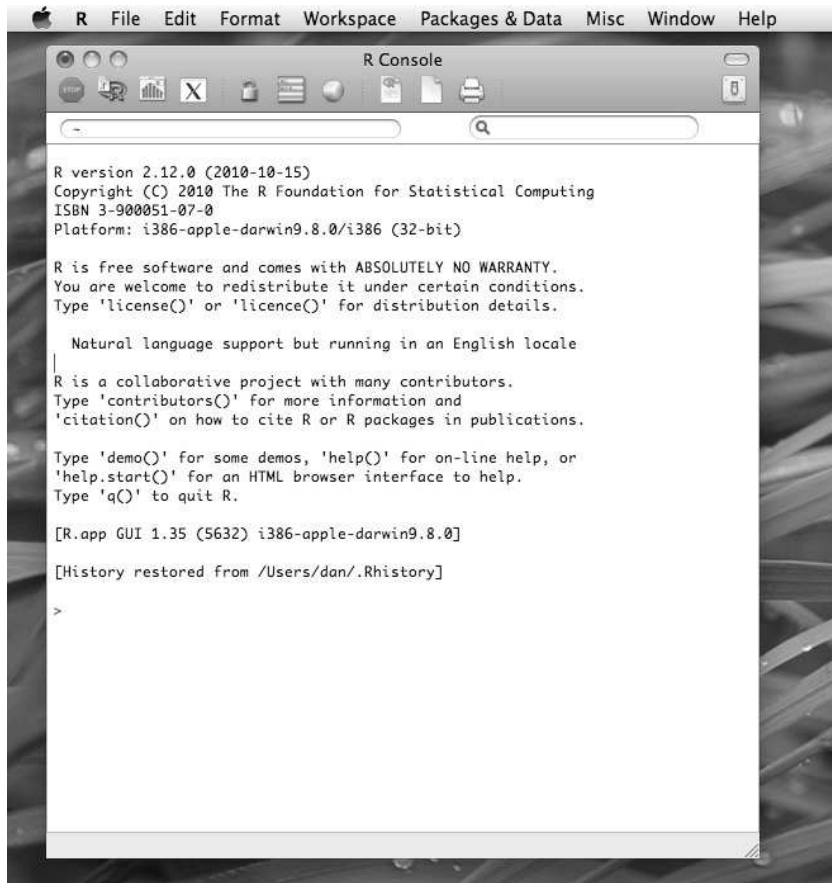


Figure 3.2: A screenshot of R running on a Mac.

(not surprisingly) called `R-2.14.0.pkg`. It's about 61MB in size, so the download can take a while on slower internet connections.

Once you've downloaded `R-2.14.0.pkg`, all you need to do is open it by double clicking on the package file. The installation should go smoothly from there: just follow all the instructions just like you usually do when you install something. Once it's finished, you'll find a file called `R.app` in the Applications folder.³ This means you now have R installed on your computer. If you double click to open R, you'll see something like the screenshot in Figure 3.2.⁴

3.1.3 Installing R on a Linux computer

If you're successfully managing to run a Linux box, regardless of what distribution, then you should find the instructions on the website easy enough. You can compile R from source yourself if you want, or install it through your package management system, which will probably have R in it. Alternatively, the

³Actually, you'll find two files, one called `R.app` and `R64.app`. The latter is the one you want for 64 bit machines. If you don't know what that means, don't worry: just ignore the pesky `R64.app` application.

⁴Tip for advanced Mac users. You can run R from the terminal if you want to. The command is just "R". It behaves like the normal desktop version, except that help documentation behaves like a "man" page instead of opening in a new window.

CRAN site has precompiled binaries for Debian, Red Hat, Suse and Ubuntu and has separate instructions for each. Once you've got R installed, you can run it from the command line just by typing `R`. Don't forget that the commands are case sensitive, so typing `r` won't work. However, if you're feeling envious of Windows and Mac users for their fancy GUIs, you can download Rstudio.

3.1.4 Downloading and installing Rstudio

At this point you're pretty much ready to go. However, there's one more step that I generally recommend that people go through: installing Rstudio. To understand what Rstudio is, you first need to understand a little bit more about R itself. The key thing to understand is the term R really refers to the underlying statistical language, and not to the "graphical user interface" (GUI) that you use to interact with R. To see this, notice that the appearance of R is very different on a Windows machine (see Figure 3.1) than on a Mac (see Figure 3.2). Some Linux users won't even have a GUI at all.

All of these are perfectly reasonable ways of using R, but they're not my favourite method, and at a practical level, it can be convenient if everyone in the class is using the same interface. To that end, my suggestion is that you also download and install Rstudio. Rstudio provides a clean, professional interface into R that I find much nicer to work with than either the Windows or Mac defaults. Like R itself, Rstudio is free software: you can find all the details on their webpage. In the meantime, you can download it here:

<http://www.rstudio.org/>

When you visit the Rstudio website, you'll probably be struck by how much cleaner and simpler it is than the CRAN website. This is probably no coincidence: the people who design and distribute the core R language itself are focused on technical stuff. And sometimes they almost seem to forget that there's an actual human user at the end. The people who design and distribute Rstudio are focused on user interface. They want to make R as usable as possible. The two websites reflect that difference.

In any case, when you click on the download button on the homepage it will ask you to choose whether you want the desktop version or the server version. At this stage you almost certainly want the desktop version. After choosing the desktop version it will take you to a page (<http://www.rstudio.org/download/desktop>) that shows several possible downloads: there's a different one for each operating system. However, the nice people at Rstudio have designed the webpage so that it automatically recommends the download that is most appropriate for your computer. Click on the appropriate link, and the Rstudio installer file will start downloading.

Once it's finished downloading, open the installer file in the usual way to install Rstudio. After it's finished installing, you can start R by opening Rstudio. To illustrate what Rstudio looks like, Figure 3.3 shows a screenshot of an R session in progress. In this screenshot, you can see that it's running on a computer with the Windows 7 operating system, but it looks almost identical no matter what operating system you have. The Mac version looks more like a Mac application (e.g., the menus are up the top and the colour scheme is grey rather than blue), but it's more or less identical. There are a few minor differences in where things are located in the menus (I'll point them out as we go along) and in the shortcut keys, because Rstudio is trying to "feel" like a proper Mac application or a proper Windows application, and this means that it has to change its behaviour a little bit depending on what computer it's running on. Even so, these differences are very small: I started out using the Mac version of Rstudio and then started using the Windows version as well in order to write these notes.⁵

The only "shortcoming" I've found with Rstudio is that – as of this writing – it's still a work in progress. The current version as I type this is 0.94, which means that it's officially still in beta testing.

⁵I don't currently have any Linux versions running, so if you're on a Linux machine you'll have to work out any differences yourself. By the time the next edition rolls around, I'll have at least one Linux distro running as well, but that's no help to you right now. I'm mentioning this more as a promissory note and an apology

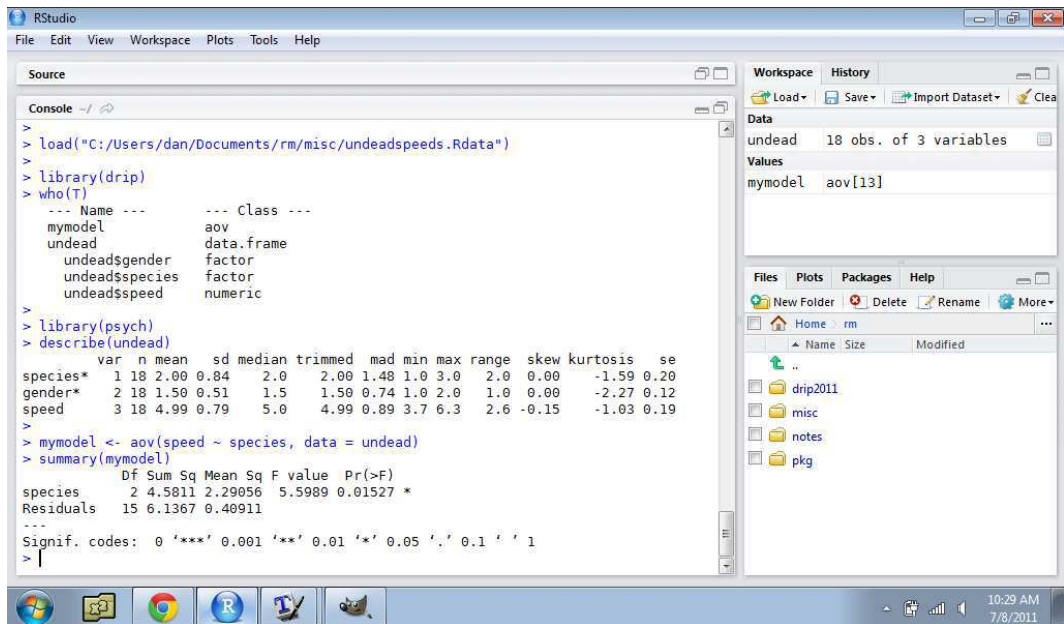


Figure 3.3: An R session in progress running through Rstudio. The operating system here is Windows 7.

Even so, the beta version of Rstudio still provides a better user experience than anything else I've tried: it really is the best option available in my opinion. The “problem” is that they keep improving it. New features keep turning up the more recent releases, so there's a good chance that by the time you read this book there will be a version out that has some really neat things that weren't in the 0.94 version.

3.1.5 Starting up R

One way or another, regardless of what operating system you're using and regardless of whether you're using Rstudio, or the default GUI, or even the command line, it's time to open R and get started. When you do that, the first thing you'll see (assuming that you're looking at the **R console**, that is) is a whole lot of text that doesn't make much sense. It should look something like this:

```
R version 2.13.1 (2011-07-08)
Copyright (C) 2011 The R Foundation for Statistical Computing
ISBN 3-900051-07-0
Platform: i386-pc-mingw32/i386 (32-bit)
```

```
R is free software and comes with ABSOLUTELY NO WARRANTY.
You are welcome to redistribute it under certain conditions.
Type 'license()' or 'licence()' for distribution details.
```

```
Natural language support but running in an English locale
```

```
R is a collaborative project with many contributors.
Type 'contributors()' for more information and
'citation()' on how to cite R or R packages in publications.
```

```
Type 'demo()' for some demos, 'help()' for on-line help, or  
'help.start()' for an HTML browser interface to help.  
Type 'q()' to quit R.
```

```
>
```

Most of this text is pretty uninteresting, and when doing real data analysis you'll never really pay much attention to it. The important part of it is this...

```
>
```

... which has a flashing cursor next to it. That's the **command prompt**. When you see this, it means that R is waiting patiently for you to do something!

3.2

Typing commands at the R console

One of the easiest things you can do with R is use it as a simple calculator, so it's a good place to start. For instance, try typing `10 + 20`, and hitting enter.⁶ When you do this, you've entered a **command**, and R will "execute" that command. What you see on screen now will be this:

```
> 10 + 20  
[1] 30
```

Not a lot of surprises in this extract. But there's a few things worth talking about, even with such a simple example. Firstly, it's important that you understand how to read the extract. In this example, what *I* typed was the `10 + 20` part. I didn't type the `>` symbol: that's just the R command prompt and isn't part of the actual command. And neither did I type the `[1] 30` part. That's what R printed out in response to my command.

Secondly, it's important to understand how the output is formatted. Obviously, the correct answer to the sum `10 + 20` is `30`, and not surprisingly R has printed that out as part of its response. But it's also printed out this `[1]` part, which probably doesn't make a lot of sense to you right now. You're going to see that a lot. I'll talk about what this means in a bit more detail later on, but for now you can think of `[1] 30` as if R were saying "the answer to the 1st question you asked is 30". That's not quite the truth, but it's close enough for now. And in any case it's not really very interesting at the moment: we only asked R to calculate one thing, so obviously there's only one answer printed on the screen. Later on this will change, and the `[1]` part will start to make a bit more sense. For now, I just don't want you to get confused or concerned by it.

3.2.1 Be very careful to avoid typos

Before we go on to talk about other types of calculations that we can do with R, there's a few other things I want to point out. The first thing is that, while R is good software, it's still software. It's pretty stupid, and because it's stupid it can't handle typos. It takes it on faith that you meant to type *exactly*

⁶Seriously. If you're in a position to do so, open up R and start typing. The simple act of typing it rather than "just reading" makes a big difference. It makes the concepts more concrete, and it ties the abstract ideas (programming and statistics) to the actual context in which you need to use them. Statistics is something you *do*, not just something you read about in a textbook.

what you did type. For example, suppose that you forgot to hit the shift key when trying to type `+`, and as a result your command ended up being `10 = 20` rather than `10 + 20`. Here's what happens:

```
> 10 = 20
Error in 10 = 20 : invalid (do_set) left-hand side to assignment
```

What's happened here is that R has attempted to interpret `10 = 20` as a command, and spits out an error message because the command doesn't make any sense to it. When a *human* looks at this, and then looks down at his or her keyboard and sees that `+` and `=` are on the same key, it's pretty obvious that the command was a typo. But R doesn't know this, so it gets upset. And, if you look at it from its perspective, this makes sense. All that R "knows" is that `10` is a legitimate number, `20` is a legitimate number, and `=` is a legitimate part of the language too. In other words, from its perspective this really does look like the user meant to type `10 = 20`, since all the individual parts of that statement are legitimate and it's too stupid to realise that this is probably a typo. Therefore, R takes it on faith that this is exactly what you meant... it only "discovers" that the command is nonsense when it tries to follow your instructions, typo and all. And then it whinges, and spits out an error.

Even more subtle is the fact that some typos won't produce errors at all, because they happen to correspond to "well-formed" R commands. For instance, suppose that not only did I forget to hit the shift key when trying to type `10 + 20`, I also managed to press the key next to one I meant do. The resulting typo would produce the command `10 - 20`. Clearly, R has no way of knowing that you meant to *add* 20 to 10, not *subtract* 20 from 10, so what happens this time is this:

```
> 10 - 20
[1] -10
```

In this case, R produces the right answer, but to the wrong question.

To some extent, I'm stating the obvious here, but it's important. The people who wrote R are smart. You, the user, are smart. But R itself is dumb. And because it's dumb, it has to be mindlessly obedient. It does *exactly* what you ask it to do. There is no equivalent to "autocorrect" in R, and for good reason. When doing advanced stuff – and even the simplest of statistics is pretty advanced in a lot of ways – it's dangerous to let a mindless automaton like R try to overrule the human user. But because of this, it's your responsibility to be careful. Always make sure you type *exactly what you mean*. When dealing with computers, it's not enough to type "approximately" the right thing. In general, you absolutely *must* be precise in what you say to R ... like all machines it is too stupid to be anything other than absurdly literal in its interpretation.

3.2.2 R is (a bit) flexible with spacing

Of course, now that I've been so uptight about the importance of always being precise, I should point out that there are some exceptions. Or, more accurately, there are some situations in which R does show a bit more flexibility than my previous description suggests. The first thing R is smart enough to do is ignore redundant spacing. What I mean by this is that, when I typed `10 + 20` before, I could equally have done this

```
> 10    + 20
[1] 30
```

or this

```
> 10+20
[1] 30
```

and I would get exactly the same answer. However, that doesn't mean that you can insert spaces in any old place. When we looked at the startup documentation in Section 3.1.5 it suggested that you could type `citation()` to get some information about how to cite R. If I do so...

```
> citation()
```

To cite R in publications use:

```
R Development Core Team (2011). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL http://www.R-project.org/.
```

BLAH BLAH BLAH

We have invested a lot of time and effort in creating R, please cite it when using it for data analysis. See also `citation("pkname")` for citing R packages.

... it tells me to cite the R manual ([R Development Core Team, 2011](#)). Obviously, the BLAH BLAH BLAH part isn't actually part of what R prints out: when you see that it means that I've choppoed out some parts of the output that I don't think are very interesting or relevant. I'll do that a lot. Anyway, getting back to my original point, let's see what happens when I try changing the spacing. If I insert spaces in between the word and the parentheses, or inside the parentheses themselves, then all is well. That is, either of these two commands

```
> citation ()  
> citation( )
```

will produce exactly the same response. However, what I can't do is insert spaces in the middle of the word. If I try to do this, R gets upset:

```
> citat ion()  
Error: unexpected symbol in "citat ion"
```

Throughout this book I'll vary the way I use spacing a little bit, just to give you a feel for the different ways in which spacing can be used. I'll try not to do it too much though, since it's generally considered to be good practice to be consistent in how you format your commands.

3.2.3 R can sometimes tell that you're not finished yet (but not often)

One more thing I should point out. If you hit enter in a situation where it's "obvious" to R that you haven't actually finished typing the command, R is just smart enough to keep waiting. For example, if you type `10 +` and then press enter, even R is smart enough to realise that you probably wanted to type in another number. So here's what happens:

```
> 10+  
+
```

and there's a blinking cursor next to the plus sign. What this means is that R is still waiting for you to finish. It "thinks" you're still typing your command, so it hasn't tried to execute it yet. In other words, this plus sign is actually another command prompt. It's different from the usual one (i.e., the > symbol) to remind you that R is going to "add" whatever you type now to what you typed last time. For example, if I then go on to type `3` and hit enter, what I get is this:

```
> 10+
+ 20
[1] 30
```

And as far as R is concerned, this is *exactly* the same as if you had typed `10 + 20`. Similarly, consider the `citation()` command that we talked about in the previous section. Suppose you hit enter after typing `citation()`. Once again, R is smart enough to realise that there must be more coming – since you need to add the `)` character – so it waits. I can even hit enter several times and it will keep waiting:

```
> citation(
+ 
+ 
+ )
```

I'll make use of this a lot in this book. A lot of the commands that we'll have to type are pretty long, and they're visually a bit easier to read if I break it up over several lines.

That being said, it's not often the case that R is smart enough to tell that there's more coming. For instance, in the same way that I can't add a space in the middle of a word, I can't hit enter in the middle of a word either. If I hit enter after typing `citat` I get an error, because R thinks I'm interested in an “object” called `citat` and can't find it:

```
> citat
Error: object 'citat' not found
```

What about if I typed `citation` and hit enter? In this case we get something very odd, something that we definitely *don't* want, at least at this stage. Here's what happens:

```
> citation
function (package = "base", lib.loc = NULL, auto = NULL)
{
  dir <- system.file(package = package, lib.loc = lib.loc)
  if (dir == "") 
    stop(gettextf("package '%s' not found", package), domain = NA)

BLAH BLAH BLAH
```

where the `BLAH BLAH BLAH` goes on for rather a long time, and you don't know enough R yet to understand what all this gibberish actually means. This incomprehensible output can be quite intimidating to novice users, and unfortunately it's very easy to forget to type the parentheses; so almost certainly you'll do this by accident. Do not panic when this happens. Simply ignore the gibberish. As you become more experienced this gibberish will start to make sense, and you'll find it quite handy to print this stuff out.⁷ But for now just try to remember to add the parentheses when typing your commands.

3.3 _____

Doing simple calculations with R

Okay, now that we've discussed some of the tedious details associated with typing R commands, let's get back to learning how to use the most powerful piece of statistical software in the world as a \$2 calculator. So far, all we know how to do is addition. Clearly, a calculator that only did addition would

⁷For advanced users: yes, as you've probably guessed, R is printing out the source code for the function.

Table 3.1: Basic arithmetic operations in R. These five operators are used very frequently throughout the text, so it's important to be familiar with them at the outset. There are others as well, which I'll discuss in Chapter 7.

operation	operator	example input	example output
addition	<code>+</code>	<code>10 + 2</code>	12
subtraction	<code>-</code>	<code>9 - 3</code>	6
multiplication	<code>*</code>	<code>5 * 5</code>	25
division	<code>/</code>	<code>10 / 3</code>	3
power	<code>^</code>	<code>5 ^ 2</code>	25

be a bit stupid, so I should tell you about how to perform other simple calculations using R. But first, some more terminology. Addition is an example of an “operation” that you can perform (specifically, an arithmetic operation), and the **operator** that performs it is `+`. To people with a programming or mathematics background, this terminology probably feels pretty natural, but to other people it might feel like I'm trying to make something very simple (addition) sound more complicated than it is (by calling it an arithmetic operation). To some extent, that's true: if addition was the only operation that we were interested in, it'd be a bit silly to introduce all this extra terminology. However, as we go along, we'll start using more and more different kinds of operations, so it's probably a good idea to get the language straight now, while we're still talking about very familiar concepts like addition!

3.3.1 Adding, subtracting, multiplying and dividing

So, now that we have the terminology, let's learn how to perform some arithmetic operations in R. To that end, Table 3.1 lists the operators that correspond to the basic arithmetic we learned in primary school: addition, subtraction, multiplication and division. As you can see, R uses fairly standard symbols to denote each of the different operations you might want to perform: addition is done using the `+` operator, subtraction is performed by the `-` operator, and so on. So if I wanted to find out what 57 times 61 is (and who wouldn't?), I can use R instead of a calculator, like so:

```
> 57 * 61
[1] 3477
```

So that's handy.

3.3.2 Taking powers

The first four operations listed in Table 3.1 are things we all learned in primary school, but they aren't the only arithmetic operations built into R. There are three other arithmetic operations that I should probably mention: taking powers, doing integer division, and calculating a modulus. Of the three, the only one that is of any real importance for the purposes of this book is taking powers, so I'll discuss that one here: the other two are discussed in Chapter 7.

For those of you who can still remember your high school maths, this should be familiar. But for some people high school maths was a long time ago, and others of us didn't listen very hard in high school. It's not complicated. As I'm sure everyone will probably remember the moment they read this, the act of multiplying a number x by itself n times is called “raising x to the n -th power”. Mathematically, this

is written as x^n . Some values of n have special names: in particular x^2 is called *x-squared*, and x^3 is called *x-cubed*. So, the 4th power of 5 is calculated like this:

$$5^4 = 5 \times 5 \times 5 \times 5$$

One way that we could calculate 5^4 in R would be to type in the complete multiplication as it is shown in the equation above. That is, we could do this

```
> 5 * 5 * 5 * 5  
[1] 625
```

but it does seem a bit tedious. It would be very annoying indeed if you wanted to calculate 5^{15} , since the command would end up being quite long. Therefore, to make our lives easier, we use the power operator instead. When we do that, our command to calculate 5^4 goes like this:

```
> 5 ^ 4  
[1] 625
```

Much easier.

3.3.3 Doing calculations in the right order

Okay. At this point, you know how to take one of the most powerful pieces of statistical software in the world, and use it as a \$2 calculator. And as a bonus, you've learned a few very basic programming concepts. That's not nothing (you could argue that you've just saved yourself \$2) but on the other hand, it's not very much either. In order to use R more effectively, we need to introduce more programming concepts.

In most situations where you would want to use a calculator, you might want to do multiple calculations. R lets you do this, just by typing in longer commands. In fact, we've already seen an example of this earlier, when I typed in `5 * 5 * 5 * 5`. However, let's try a slightly different example:

```
> 1 + 2 * 4  
[1] 9
```

Clearly, this isn't a problem for R either. However, it's worth stopping for a second, and thinking about what R just did. Clearly, since it gave us an answer of 9 it must have multiplied `2 * 4` (to get an interim answer of 8) and then added 1 to that. But, suppose it had decided to just go from left to right: if R had decided instead to add `1+2` (to get an interim answer of 3) and then multiplied by 4, it would have come up with an answer of 12.

To answer this, you need to know the **order of operations** that R uses. If you remember back to your high school maths classes, it's actually the same order that you got taught when you were at school: the "BEDMAS" order.⁸ That is, first calculate things inside **Brackets** (), then calculate **Exponents** ^, then **Division** / and **Multiplication** *, then **Addition** + and **Subtraction** -. So, to continue the example above, if we want to force R to calculate the `1+2` part before the multiplication, all we would have to do is enclose it in brackets:

```
> (1 + 2) * 4  
[1] 12
```

⁸For advanced users: if you want a table showing the complete order of operator precedence in R, type `?Syntax`. I haven't included it in this book since there are quite a few different operators, and we don't need that much detail. Besides, in practice most people seem to figure it out from seeing examples: until writing this book I never looked at the formal statement of operator precedence for any language I ever coded in, and never ran into any difficulties.

This is a fairly useful thing to be able to do. The only other thing I should point out about order of operations is what to expect when you have two operations that have the same priority: that is, how does R resolve ties? For instance, multiplication and division are actually the same priority, but what should we expect when we give R a problem like `4 / 2 * 3` to solve? If it evaluates the multiplication first and then the division, it would calculate a value of two-thirds. But if it evaluates the division first it calculates a value of 6. The answer, in this case, is that R goes from *left to right*, so in this case the division step would come first:

```
> 4 / 2 * 3  
[1] 6
```

All of the above being said, it's helpful to remember that *brackets always come first*. So, if you're ever unsure about what order R will do things in, an easy solution is to enclose the thing *you* want it to do first in brackets. There's nothing stopping you from typing `(4 / 2) * 3`. By enclosing the division in brackets we make it clear which thing is supposed to happen first. In this instance you wouldn't have needed to, since R would have done the division first anyway, but when you're first starting out it's better to make sure R does what you want!

3.4

Storing a number as a variable

One of the most important things to be able to do in R (or any programming language, for that matter) is to store information in **variables**. Variables in R aren't exactly the same thing as the variables we talked about in the last chapter on research methods, but they are similar. At a conceptual level you can think of a variable as *label* for a certain piece of information, or even several different pieces of information. When doing statistical analysis in R all of your data (the variables you measured in your study) will be stored as variables in R, but as well see later in the book you'll find that you end up creating variables for other things too. However, before we delve into all the messy details of data sets and statistical analysis, let's look at the very basics for how we create variables and work with them.

3.4.1 Variable assignment using `<-` and `->`

Since we've been working with numbers so far, let's start by creating variables to store our numbers. And since most people like concrete examples, let's invent one. Suppose I'm trying to calculate how much money I'm going to make from this book. There's several different numbers I might want to store. Firstly, I need to figure out how many copies I'll sell. This isn't exactly *Harry Potter*, so let's assume I'm only going to sell one copy per student in my class. That's 350 sales, so let's create a variable called **sales**. What I want to do is assign a **value** to my variable **sales**, and that value should be **350**. We do this by using the **assignment operator**, which is `<-`. Here's how we do it:

```
> sales <- 350
```

When you hit enter, R doesn't print out any output.⁹ It just gives you another command prompt. However, behind the scenes R has created a variable called **sales** and given it a value of **350**. You can

⁹If you are using Rstudio, and the “workspace” panel is visible when you typed the command, then you probably saw something happening there. That's to be expected, and is quite helpful. However, there's two things to note here (1) I haven't yet explained what that panel does, so for now just ignore it, and (2) this is one of the helpful things Rstudio does, not a part of R itself.

check that this has happened by asking R to print the variable on screen. And the simplest way to do *that* is to type the name of the variable and hit enter¹⁰

```
> sales  
[1] 350
```

So that's nice to know. Anytime you can't remember what R has got stored in a particular variable, you can just type the name of the variable and hit enter.

Okay, so now we know how to assign variables. Actually, there's a bit more you should know. Firstly, one of the curious features of R is that there are several different ways of making assignments. In addition to the `<-` operator, we can also use `->` and `=`, and it's pretty important to understand the differences between them.¹¹ Let's start by considering `->`, since that's the easy one (we'll discuss the use of `=` in Section 3.5.1). As you might expect from just looking at the symbol, it's almost identical to `<-`. It's just that the arrow (i.e., the assignment) goes from left to right. So if I wanted to define my `sales` variable using `->`, I would write it like this:

```
> 350 -> sales
```

This has the same effect: and it *still* means that I'm only going to sell 350 copies. Sigh. Apart from this superficial difference, `<-` and `->` are identical. In fact, as far as R is concerned, they're actually the same operator, just in a "left form" and a "right form".¹²

3.4.2 Doing calculations using variables

Okay, let's get back to my original story. In my quest to become rich, I've written this textbook. To figure out how good a strategy is, I've started creating some variables in R. In addition to defining a `sales` variable that counts the number of copies I'm going to sell, I can also create a variable called `royalty`, indicating how much money I get per copy. Let's say that my royalties are about \$7 per book:

```
> sales <- 350  
> royalty <- 7
```

The nice thing about variables (in fact, the whole point of having variables) is that we can do anything with a variable that we ought to be able to do with the information that it stores. That is, since R allows me to multiply `350` by `7`

```
> 350 * 7  
[1] 2450
```

it also allows me to multiply `sales` by `royalty`

```
> sales * royalty  
[1] 2450
```

As far as R is concerned, the `sales * royalty` command is the same as the `350 * 7` command. Not surprisingly, I can assign the output of this calculation to a new variable, which I'll call `revenue`. And when we do this, the new variable `revenue` gets the value `2450`. So let's do that, and then get R to print out the value of `revenue` so that we can verify that it's done what we asked:

¹⁰As we'll discuss later, by doing this we are implicitly using the `print()` function.

¹¹Actually, in keeping with the R tradition of providing you with a billion different screwdrivers (even when you're actually looking for a hammer) these aren't the only options. There's also the `assign()` function, and the `<<-` and `->>` operators. However, we won't be using these at all in this book.

¹²A quick reminder: when using operators like `<-` and `->` that span multiple characters, you can't insert spaces in the middle. That is, if you type `- >` or `< -`, R will interpret your command the wrong way. And I will cry.

```
> revenue <- sales * royalty
> revenue
[1] 2450
```

That's fairly straightforward. A slightly more subtle thing we can do is reassign the value of my variable, based on its current value. For instance, suppose that one of my students (no doubt under the influence of psychotropic drugs) loves the book so much that he or she donates me an extra \$550. The simplest way to capture this is by a command like this:

```
> revenue <- revenue + 550
> revenue
[1] 3000
```

In this calculation, R has taken the old value of `revenue` (i.e., 2450) and added 550 to that value, producing a value of 3000. This new value is assigned to the `revenue` variable, overwriting its previous value. In any case, we now know that I'm expecting to make \$3000 off this. Pretty sweet, I thinks to myself. Or at least, that's what I thinks until I do a few more calculation and work out what the implied hourly wage I'm making off this looks like.

3.4.3 Rules and conventions for naming variables

In the examples that we've seen so far, my variable names (`sales` and `revenue`) have just been English-language words written using lowercase letters. However, R allows a lot more flexibility when it comes to naming your variables, as the following list of rules¹³ illustrates:

- Variable names can only use the upper case alphabetic characters `A-Z` as well as the lower case characters `a-z`. You can also include numeric characters `0-9` in the variable name, as well as the period `.` or underscore `_` character. In other words, you can use `SaL.e_s` as a variable name (though I can't think why you would want to), but you can't use `Sales?`.
- Variable names cannot include spaces: therefore `my sales` is not a valid name, but `my.sales` is.
- Variable names are case sensitive: that is, `Sales` and `sales` are *different* variable names.
- Variable names must start with a letter or a period. You can't use something like `_sales` or `1sales` as a variable name. You can use `.sales` as a variable name if you want, but it's not usually a good idea. By convention, variables starting with a `.` are used for special purposes, so you should avoid doing so.
- Variable names cannot be one of the reserved keywords. These are special names that R needs to keep "safe" from us mere users, so you can't use them as the names of variables. The keywords are: `if`, `else`, `repeat`, `while`, `function`, `for`, `in`, `next`, `break`, `TRUE`, `FALSE`, `NULL`, `Inf`, `NaN`, `NA`, `NA_integer_`, `NA_real_`, `NA_complex_`, and finally, `NA_character_`. Don't feel especially obliged to memorise these: if you make a mistake and try to use one of the keywords as a variable name, R will complain about it like the whiny little automaton it is.

In addition to those rules that R enforces, there are some informal conventions that people tend to follow when naming variables. One of them you've already seen: i.e., don't use variables that start with a period. But there are several others. You aren't obliged to follow these conventions, and there are many situations in which it's advisable to ignore them, but it's generally a good idea to follow them when you can:

¹³Actually, you can override any of these rules if you want to, any quite easily. All you have to do is add quote marks or backticks around your non-standard variable name. For instance '`'my sales'` `<- 350` would work just fine, but it's almost never a good idea to do this.

- Use informative variable names. As a general rule, using meaningful names like `sales` and `revenue` is preferred over arbitrary ones like `variable1` and `variable2`. Otherwise it's very hard to remember what the contents of different variables are, and it becomes hard to understand what your commands actually do.
- Use short variable names. Typing is a pain and no-one likes doing it. So we much prefer to use a name like `sales` over a name like `sales.for.this.book.that.you.are.reading`. Obviously there's a bit of a tension between using informative names (which tend to be long) and using short names (which tend to be meaningless), so use a bit of common sense when trading off these two conventions.
- Use one of the conventional naming styles for multi-word variable names. Suppose I want to name a variable that stores "my new salary". Obviously I can't include spaces in the variable name, so how should I do this? There are three different conventions that you sometimes see R users employing. Firstly, you can separate the words using periods, which would give you `my.new.salary` as the variable name. Alternatively, you could separate words using underscores, as in `my_new_salary`. Finally, you could use capital letters at the beginning of each word (except the first one), which gives you `myNewSalary` as the variable name. I don't think there's any strong reason to prefer one over the other,¹⁴ but it's important to be consistent.

3.5

Using functions to do calculations

The symbols `+`, `-`, `*` and so on are examples of operators. As we've seen, you can do quite a lot of calculations just by using these operators. However, in order to do more advanced calculations (and later on, to do actual statistics), you're going to need to start using `functions`.¹⁵ I'll talk in more detail about functions and how they work in Section 8.4, but for now let's just dive in and use a few. To get started, suppose I wanted to take the square root of 225. The square root, in case your high school maths is a bit rusty, is just the opposite of squaring a number. So, for instance, since "5 squared is 25" I can say that "5 is the square root of 25". The usual notation for this is

$$\sqrt{25} = 5$$

though sometimes you'll also see it written like this $25^{0.5} = 5$. This second way of writing it is kind of useful to "remind" you of the mathematical fact that "square root of x " is actually the same as "raising x to the power of 0.5". Personally, I've never found this to be terribly meaningful psychologically, though I have to admit it's quite convenient mathematically. Anyway, it's not important. What is important is that you remember what a square root is, since we're going to need it later on.

To calculate the square root of 25, I can do it in my head pretty easily, since I memorised my multiplication tables when I was a kid. It gets harder when the numbers get bigger, and pretty much impossible if they're not whole numbers. This is where something like R comes in very handy. Let's say I wanted to calculate $\sqrt{225}$, the square root of 225. There's two ways I could do this using R. Firstly, since the square root of 225 is the same thing as raising 225 to the power of 0.5, I could use the power operator `^`, just like we did earlier:

¹⁴For very advanced users: there is one exception to this. If you're naming a function, don't use `.` in the name unless you are intending to make use of the S3 object oriented programming system in R. If you don't know what S3 is, then you definitely don't want to be using it! For function naming, there's been a trend among R users to prefer `myFunctionName`.

¹⁵A side note for students with a programming background. Technically speaking, operators *are* functions in R: the addition operator `+` is actually a convenient way of calling the addition function `'+'()`. Thus `10+20` is equivalent to the function call `'+'(20, 30)`. Not surprisingly, no-one ever uses this version. Because that would be stupid.

```
> 225 ^ 0.5  
[1] 15
```

However, there's a second way that we can do this, since R also provides a **square root function**, `sqrt()`. To calculate the square root of 255 using this function, what I do is insert the number `225` in the parentheses. That is, the command I type is this:

```
> sqrt( 225 )  
[1] 15
```

and as you might expect from our previous discussion, the spaces in between the parentheses are purely cosmetic. I could have typed `sqrt(225)` or `sqrt(225)` and gotten the same result. When we use a function to do something, we generally refer to this as **calling** the function, and the values that we type into the function (there can be more than one) are referred to as the **arguments** of that function.

Obviously, the `sqrt()` function doesn't really give us any new functionality, since we already knew how to do square root calculations by using the power operator `^`, though I do think it looks nicer when we use `sqrt()`. However, there are lots of other functions in R: in fact, almost everything of interest that I'll talk about in this book is an R function of some kind. For example, one function that we will need to use in this book is the **absolute value function**. Compared to the square root function, it's extremely simple: it just converts negative numbers to positive numbers, and leaves positive numbers alone. Mathematically, the absolute value of x is written $|x|$ or sometimes $\text{abs}(x)$. Calculating absolute values in R is pretty easy, since R provides the `abs()` function that you can use for this purpose. When you feed it a positive number...

```
> abs( 21 )  
[1] 21
```

the absolute value function does nothing to it at all. But when you feed it a negative number, it spits out the positive version of the same number, like this:

```
> abs( -13 )  
[1] 13
```

In all honesty, there's nothing that the absolute value function does that you couldn't do just by looking at the number and erasing the minus sign if there is one. However, there's a few places later in the book where we have to use absolute values, so I thought it might be a good idea to explain the meaning of the term early on.

Before moving on, it's worth noting that – in the same way that R allows us to put multiple operations together into a longer command, like `1 + 2*4` for instance – it also lets us put functions together and even combine functions with operators if we so desire. For example, the following is a perfectly legitimate command:

```
> sqrt( 1 + abs(-8) )  
[1] 3
```

When R executes this command, starts out by calculating the value of `abs(-8)`, which produces an intermediate value of `8`. Having done so, the command simplifies to `sqrt(1 + 8)`. To solve the square root¹⁶ it first needs to add `1 + 8` to get `9`, at which point it evaluates `sqrt(9)`, and so it finally outputs a value of `3`.

¹⁶A note for the mathematically inclined: R does support complex numbers, but unless you explicitly specify that you want them it assumes all calculations must be real valued. By default, the square root of a negative number is treated as undefined: `sqrt(-9)` will produce `NaN` (not a number) as its output. To get complex numbers, you would type `sqrt(-9+0i)` and R would now return `0+3i`. However, since we won't have any need for complex numbers in this book, I won't refer to them again.

3.5.1 Function arguments, their names and their defaults

There's two more fairly important things that you need to understand about how functions work in R, and that's the use of "named" arguments, and default values" for arguments. Not surprisingly, that's not to say that this is the last we'll hear about how functions work, but they are the last things we desperately need to discuss in order to get you started. To understand what these two concepts are all about, I'll introduce another function. The `round()` function can be used to round some value to the nearest whole number. For example, I could type this:

```
> round( 3.1415 )
[1] 3
```

Pretty straightforward, really. However, suppose I only wanted to round it to two decimal places: that is, I want to get `3.14` as the output. The `round()` function supports this, by allowing you to input a second argument to the function that specifies the number of decimal places that you want to round the number to. In other words, I could do this:

```
> round( 3.14165, 2 )
[1] 3.14
```

What's happening here is that I've specified *two* arguments: the first argument is the number that needs to be rounded (i.e., `3.1415`), the second argument is the number of decimal places that it should be rounded to (i.e., `2`), and the two arguments are separated by a comma. In this simple example, it's quite easy to remember which one argument comes first and which one comes second, but for more complicated functions this is not easy. Fortunately, most R functions make use of **argument names**. For the `round()` function, for example the number that needs to be rounded is specified using the `x` argument, and the number of decimal points that you want it rounded to is specified using the `digits` argument. Because we have these names available to us, we can specify the arguments to the function by name. We do so like this:

```
> round( x = 3.1415, digits = 2 )
[1] 3.14
```

Notice that this is kind of similar in spirit to variable assignment (Section 3.4), except that I used `=` here, rather than `<-`. In both cases we're specifying specific values to be associated with a label. However, there are some differences between what I was doing earlier on when creating variables, and what I'm doing here when specifying arguments, and so as a consequence it's important that you use `=` in this context.

As you can see, specifying the arguments by name involves a lot more typing, but it's also a lot easier to read. Because of this, the commands in this book will usually specify arguments by name,¹⁷ since that makes it clearer to you what I'm doing. However, one important thing to note is that when specifying the arguments using their names, it doesn't matter what order you type them in. But if you don't use the argument names, then you have to input the arguments in the correct order. In other words, these three commands all produce the same output...

```
> round( 3.14165, 2 )
> round( x = 3.1415, digits = 2 )
> round( digits = 2, x = 3.1415 )
```

¹⁷The two functions discussed previously, `sqrt()` and `abs()`, both only have a single argument, `x`. So I could have typed something like `sqrt(x = 225)` or `abs(x = -13)` earlier. The fact that all these functions use `x` as the name of the argument that corresponds the "main" variable that you're working with is no coincidence. That's a fairly widely used convention. Quite often, the writers of R functions will try to use conventional names like this to make your life easier. Or at least that's the theory. In practice it doesn't always work as well as you'd hope.

but this one does not...

```
> round( 2, 3.14165 )
```

How do you find out what the correct order is? There's a few different ways, but the easiest one is to look at the help documentation for the function (see Section 4.12). However, if you're ever unsure, it's probably best to actually type in the argument name.

Okay, so that's the first thing I said you'd need to know: argument names. The second thing you need to know about is default values. Notice that the first time I called the `round()` function I didn't actually specify the `digits` argument at all, and yet R somehow knew that this meant it should round to the nearest whole number. How did that happen? The answer is that the `digits` argument has a **default value** of `0`, meaning that if you decide not to specify a value for `digits` then R will act as if you had typed `digits = 0`. This is quite handy: the vast majority of the time when you want to round a number you want to round it to the nearest whole number, and it would be pretty annoying to have to specify the `digits` argument every single time. On the other hand, sometimes you actually do want to round to something other than the nearest whole number, and it would be even more annoying if R didn't allow this! Thus, by having `digits = 0` as the default value, we get the best of both worlds.

3.6

Storing many numbers as a vector

At this point we've covered functions in enough detail to get us safely through the next couple of chapters (with one small exception: see Section 4.11), so let's return to our discussion of variables. When I introduced variables in Section 3.4 I showed you how we can use variables to store a single number. In this section, we'll extend this idea and look at how to store multiple numbers within the one variable. In R the name for a variable that can store multiple values is a **vector**. So let's create one.

3.6.1 Creating a vector

Let's stick to my silly "get rich quick by textbook writing" example. Suppose the textbook company sends me sales data on a monthly basis. Since my class start in late February, we might expect most of the sales to occur towards the start of the year. Let's suppose that I have 100 sales in February, 200 sales in March and 50 sales in April, and no other sales for the rest of the year. What I would like to do is have a variable – let's call it `sales.by.month` – that stores all this sales data. The first number stored should be `0` since I had no sales in January, the second should be `100`, and so on. The simplest way to do this in R is to use the `combine` function, `c()`. To do so, all we have to do is type all the numbers you want to store in a comma separated list, like this:¹⁸

```
> sales.by.month <- c(0, 100, 200, 50, 0, 0, 0, 0, 0, 0, 0, 0)
> sales.by.month
[1] 0 100 200 50 0 0 0 0 0 0 0 0
```

To use the correct terminology here, we have a single variable here called `sales.by.month`: this variable is a vector that consists of 12 **elements**.

¹⁸Notice that I didn't specify any argument names here. The `c()` function is one of those cases where we don't use names. We just type all the numbers, and R just dumps them all in a single variable.

3.6.2 A handy digression

Now that we've learned how to put information into a vector, the next thing to understand is how to pull that information back out again. However, before I do so it's worth taking a slight detour. If you've been following along, typing all the commands into R yourself, it's possible that the output that you saw when we printed out the `sales.by.month` vector was slightly different to what I showed above. This would have happened if the window (or the Rstudio panel) that contains the R console is really, really narrow. If that were the case, you might have seen output that looks something like this:

```
> sales.by.month  
[1] 0 100 200 50 0 0 0 0  
[9] 0 0 0 0
```

Because there wasn't much room on the screen, R has printed out the results over two lines. But that's not the important thing to notice. The important point is that the first line has a `[1]` in front of it, whereas the second line starts with `[9]`. It's pretty clear what's happening here. For the first row, R has printed out the 1st element through to the 8th element, so it starts that row with a `[1]`. For the second row, R has printed out the 9th element of the vector through to the 12th one, and so it begins that row with a `[9]` so that you can tell where it's up to at a glance. It might seem a bit odd to you that R does this, but in some ways it's a kindness, especially when dealing with larger data sets!

3.6.3 Getting information out of vectors

To get back to the main story, let's consider the problem of how to get information out of a vector. At this point, you might have a sneaking suspicion that the answer has something to do with the `[1]` and `[9]` things that R has been printing out. And of course you are correct. Suppose I want to pull out the February sales data only. February is the second month of the year, so let's try this:

```
> sales.by.month[2]  
[1] 100
```

Yep, that's the February sales all right. But there's a subtle detail to be aware of here: notice that R outputs `[1] 100`, *not* `[2] 100`. This is because R is being extremely literal. When we typed in `sales.by.month[2]`, we asked R to find exactly *one* thing, and that one thing happens to be the second element of our `sales.by.month` vector. So, when it outputs `[1] 100` what R is saying is that the first number *that we just asked for* is 100. This behaviour makes more sense when you realise that we can use this trick to create new variables. For example, I could create a `february.sales` variable like this:

```
> february.sales <- sales.by.month[2]  
> february.sales  
[1] 100
```

Obviously, the new variable `february.sales` should only have one element and so when I print it out this new variable, the R output begins with a `[1]` because `100` is the value of the first (and only) element of `february.sales`. The fact that this also happens to be the value of the second element of `sales.by.month` is irrelevant. We'll pick this topic up again shortly (Section 3.9).

3.6.4 Altering the elements of a vector

Sometimes you'll want to change the values stored in a vector. Imagine my surprise when the publisher rings me up to tell me that the sales data for May are wrong. There were actually an additional 25 books

sold in May, but there was an error or something so they hadn't told me about it. How can I fix my `sales.by.month` variable? One possibility would be to assign the whole vector again from the beginning, using `c()`. But that's a lot of typing. Also, it's a little wasteful: why should R have to redefine the sales figures for all 12 months, when only the 5th one is wrong? Fortunately, we can tell R to change only the 5th element, using this trick:

```
> sales.by.month[5] <- 25
> sales.by.month
[1] 0 100 200 50 25 0 0 0 0 0 0
```

Another way to edit variables is to use the `edit()` or `fix()` functions. I won't discuss them in detail in this book, mainly because they're basically methods for visually editing variables in a spreadsheet-like format, and they're not currently very well supported in Rstudio (this is, in my opinion, one of the biggest shortcomings of Rstudio, though I still think it's the best front end for R available at the moment). They're not *completely* unsupported: if you click on the entry for `sales.by.month` in Rstudio's workspace panel, it will run the equivalent of a `fix(sales.by.month)` command. It opens up a little window that contains the actual R command that you would use to define the variable (i.e., it reads `c(0, 100, 200, 50, 0, 0, 0, 0, 0, 0, 0, 0)`) and you can manually edit that command. If you then click on the "Save" button R will redefine the `sales.by.month` variable for you. This is kind of neat, but it only works for some kinds of variables (e.g., it won't work on data frames, which you'll use a lot). The Rstudio team have indicated on their website that they intend to introduce some richer functionality here in future versions, so this is a bit of a "wait and see" kind of proposition.

3.6.5 Useful things to know about vectors

Before moving on, I want to mention a couple of other things about vectors. Firstly, you often find yourself wanting to know how many elements there are in a vector (usually because you've forgotten). You can use the `length()` function to do this. It's quite straightforward:

```
> length( x = sales.by.month )
[1] 12
```

Secondly, you often want to alter all of the elements of a vector at once. For instance, suppose I wanted to figure out how much money I made in each month. Since I'm earning an exciting \$7 per book (no seriously, that's actually pretty close to the true number), what I want to do is multiply each element in the `sales.by.month` vector by 7. R makes this pretty easy, as the following example shows:

```
> sales.by.month * 7
[1] 0 700 1400 350 0 0 0 0 0 0 0 0
```

In other words, when you multiply a vector by a single number, all elements in the vector get multiplied. The same is true for addition, subtraction, division and taking powers. So that's neat. On the other hand, suppose I wanted to know how much money I was making per day, rather than per month. Since not every month has the same number of days, I need to do something slightly different. Firstly, I'll create two new vectors:

```
> days.per.month <- c(31, 28, 31, 30, 31, 30, 31, 31, 30, 31, 30, 31)
> profit <- sales.by.month * 7
```

Obviously, the `profit` variable is the same one we created earlier, and the `days.per.month` variable is pretty straightforward. What I want to do is divide every element of `profit` by the *corresponding* element of `days.per.month`. Again, R makes this pretty easy:

```
> profit / days.per.month
[1] 0.00000 25.00000 45.16129 11.66667 0.00000 0.00000 0.00000 0.00000 0.00000
[10] 0.00000 0.00000 0.00000
```

I still don't like all those zeros, but that's not what matters here. Notice that the second element of the output is 25, because R has divided the second element of `profit` (i.e. 700) by the second element of `days.per.month` (i.e. 28). Similarly, the third element of the output is equal to 1400 divided by 31, and so on. We'll talk more about calculations involving vectors later on (and in particular a thing called the "recycling rule"; Section 7.12.2), but that's enough detail for now.

3.7

Storing text data

A lot of the time your data will be numeric in nature, but not always. Sometimes your data really needs to be described using text, not using numbers. To address this, we need to consider the situation where our variables store text. To create a variable that stores the word "hello", we can type this:

```
> greeting <- "hello"
> greeting
[1] "hello"
```

When interpreting this, it's important to recognise that the quote marks here *aren't* part of the string itself. They're just something that we use to make sure that R knows to treat the characters that they enclose as a piece of text data, known as a **character string**. In other words, R treats "`hello`" as a string containing the word "hello"; but if I had typed `hello` instead, R would go looking for a variable by that name! You can also use '`hello`' to specify a character string.

Okay, so that's how we store the text. Next, it's important to recognise that when we do this, R stores the entire word "`hello`" as a *single* element: our `greeting` variable is not a vector of five different letters. Rather, it has only the one element, and that element corresponds to the entire character string "`hello`". To illustrate this, if I actually ask R to find the first element of `greeting`, it prints the whole string:

```
> greeting[1]
[1] "hello"
```

Of course, there's no reason why I can't create a vector of character strings. For instance, if we were to continue with the example of my attempts to look at the monthly sales data for my book, one variable I might want would include the names of all 12 `months`.¹⁹ To do so, I could type in a command like this

```
> months <- c("January", "February", "March", "April", "May", "June",
+           "July", "August", "September", "October", "November", "December")
> months
[1] "January"   "February"  "March"      "April"       "May"        "June"
[7] "July"       "August"    "September" "October"    "November"   "December"
```

This is a **character vector** containing 12 elements, each of which is the name of a month. So if I wanted R to tell me the name of the fourth month, all I would do is this:

```
> months[4]
[1] "April"
```

¹⁹Though actually there's no real need to do this, since R has an inbuilt variable called `month.name` that you can use for this purpose.

3.7.1 Working with text

Working with text data is somewhat more complicated than working with numeric data, and I discuss some of the basic ideas in Section 7.8, but for purposes of the current chapter we only need this bare bones sketch. The only other thing I want to do before moving on is show you an example of a function that can be applied to text data. So far, most of the functions that we have seen (i.e., `sqrt()`, `abs()` and `round()`) only make sense when applied to numeric data (e.g., you can't calculate the square root of "hello"), and we've seen one function that can be applied to pretty much any variable or vector (i.e., `length()`). So it might be nice to see an example of a function that can be applied to text.

The function I'm going to introduce you to is called `nchar()`, and what it does is count the number of individual characters that make up a string. Recall earlier that when we tried to calculate the `length()` of our `greeting` variable it returned a value of 1: the `greeting` variable contains only the one string, which happens to be "`hello`". But what if I want to know how many letters there are in the word? Sure, I could *count* them, but that's boring, and more to the point it's a terrible strategy if what I wanted to know was the number of letters in *War and Peace*. That's where the `nchar()` function is helpful:

```
> nchar( x = greeting )
[1] 5
```

That makes sense, since there are in fact 5 letters in the string "`hello`". Better yet, you can apply `nchar()` to whole vectors. So, for instance, if I want R to tell me how many letters there are in the names of each of the 12 months, I can do this:

```
> nchar( x = months )
[1] 7 8 5 5 3 4 4 6 9 7 8 8
```

So that's nice to know. The `nchar()` function can do a bit more than this, and there's a lot of other functions that you can do to extract more information from text or do all sorts of fancy things. However, the goal here is not to teach any of that! The goal right now is just to see an example of a function that actually does work when applied to text.

3.8

Storing “true or false” data

Time to move onto a third kind of data. A key concept in that a lot of R relies on is the idea of a **logical value**. A logical value is an assertion about whether something is true or false. This is implemented in R in a pretty straightforward way. There are two logical values, namely `TRUE` and `FALSE`. Despite the simplicity, a logical values are very useful things. Let's see how they work.

3.8.1 Assessing mathematical truths

In George Orwell's classic book *1984*, one of the slogans used by the totalitarian Party was “two plus two equals five”, the idea being that the political domination of human freedom becomes complete when it is possible to subvert even the most basic of mathematical truths. It's a terrifying thought, especially when the protagonist Winston Smith finally breaks down under torture and agrees to the proposition. “Man is infinitely malleable”, the book says. I'm pretty sure that this isn't true of humans²⁰ but it's

²⁰I offer up my teenage attempts to be “cool” as evidence that some things just can't be done.

definitely not true of R. R is not infinitely malleable. It has rather firm opinions on the topic of what is and isn't true, at least as regards basic mathematics. If I ask it to calculate `2 + 2`, it always gives the same answer, and it's not bloody 5:

```
> 2 + 2  
[1] 4
```

Of course, so far R is just doing the calculations. I haven't asked it to explicitly assert that $2 + 2 = 4$ is a true statement. If I want R to make an explicit judgement, I can use a command like this:

```
> 2 + 2 == 4  
[1] TRUE
```

What I've done here is use the **equality operator**, `==`, to force R to make a “true or false” judgement.²¹ Okay, let's see what R thinks of the Party slogan:

```
> 2+2 == 5  
[1] FALSE
```

Booyah! Freedom and ponies for all! Or something like that. Anyway, it's worth having a look at what happens if I try to *force* R to believe that two plus two is five by making an assignment statement like `2 + 2 = 5` or `2 + 2 <- 5`. When I do this, here's what happens:

```
> 2 + 2 = 5  
Error in 2 + 2 = 5 : target of assignment expands to non-language object
```

R doesn't like this very much. It recognises that `2 + 2` is *not* a variable (that's what the “non-language object” part is saying), and it won't let you try to “reassign” it. While R is pretty flexible, and actually does let you do some quite remarkable things to redefine parts of R itself, there are just some basic, primitive truths that it refuses to give up. It won't change the laws of addition, and it won't change the definition of the number `2`.

That's probably for the best.

3.8.2 Logical operations

So now we've seen logical operations at work, but so far we've only seen the simplest possible example. You probably won't be surprised to discover that we can combine logical operations with other operations and functions in a more complicated way, like this:

```
> 3*3 + 4*4 == 5*5  
[1] TRUE
```

or this

```
> sqrt( 25 ) == 5  
[1] TRUE
```

²¹Note that this is a very different operator to the assignment operator `=` that I talked about in Section 3.4. A common typo that people make when trying to write logical commands in R (or other languages, since the “`=` versus `==`” distinction is important in most programming languages) is to accidentally type `=` when you really mean `==`. Be especially cautious with this – I've been programming in various languages since I was a teenager, and I *still* screw this up a lot. Hm. I think I see why I wasn't cool as a teenager. And why I'm still not cool.

Table 3.2: Some logical operators. Technically I should be calling these “binary relational operators”, but quite frankly I don’t want to. It’s my book so no-one can make me.

operation	operator	example input	answer
less than	<	2 < 3	TRUE
less than or equal to	<=	2 <= 2	TRUE
greater than	>	2 > 3	FALSE
greater than or equal to	>=	2 >= 2	TRUE
equal to	==	2 == 3	FALSE
not equal to	!=	2 != 3	TRUE

Not only that, but as Table 3.2 illustrates, there are several other logical operators that you can use, corresponding to some basic mathematical concepts. Hopefully these are all pretty self-explanatory: for example, the **less than** operator `<` checks to see if the number on the left is less than the number on the right. If it’s less, then R returns an answer of `TRUE`:

```
> 99 < 100
[1] TRUE
```

but if the two numbers are equal, or if the one on the right is larger, then R returns an answer of `FALSE`, as the following two examples illustrate:

```
> 100 < 100
[1] FALSE
> 100 < 99
[1] FALSE
```

In contrast, the **less than or equal to** operator `<=` will do exactly what it says. It returns a value of `TRUE` if the number of the left hand side is less than or equal to the number on the right hand side. So if we repeat the previous two examples using `<=`, here’s what we get:

```
> 100 <= 100
[1] TRUE
> 100 <= 99
[1] FALSE
```

And at this point I hope it’s pretty obvious what the **greater than** operator `>` and the **greater than or equal to** operator `>=` do! Next on the list of logical operators is the **not equal to** operator `!=` which – as with all the others – does what it says it does. It returns a value of `TRUE` when things on either side are not identical to each other. Therefore, since $2 + 2 \neq 5$, we get:

```
> 2 + 2 != 5
[1] TRUE
```

We’re not quite done yet. There are three more logical operations that are worth knowing about, listed in Table 3.3. These are the **not** operator `!`, the **and** operator `&`, and the **or** operator `|`. Like the other logical operators, their behaviour is more or less exactly what you’d expect given their names. For instance, if I ask you to assess the claim that “either $2 + 2 = 4$ or $2 + 2 = 5$ ” you’d say that it’s true. Since it’s an “either-or” statement, all we need is for one of the two parts to be true. That’s what the `|` operator does:

Table 3.3: Some more logical operators.

operation	operator	example input	answer
not	!	<code>!(1==1)</code>	FALSE
or		<code>(1==1) (2==3)</code>	TRUE
and	&	<code>(1==1) & (2==3)</code>	FALSE

```
> (2+2 == 4) | (2+2 == 5)
[1] TRUE
```

On the other hand, if I ask you to assess the claim that “both $2 + 2 = 4$ and $2 + 2 = 5$ ” you’d say that it’s false. Since this is an *and* statement we need both parts to be true. And that’s what the `&` operator does:

```
> (2+2 == 4) & (2+2 == 5)
[1] FALSE
```

Finally, there’s the *not* operator, which is simple but annoying to describe in English. If I ask you to assess my claim that “it is not true that $2 + 2 = 5$ ” then you would say that my claim is true; because my claim is that “ $2 + 2 = 5$ is false”. And I’m right. If we write this as an R command we get this:

```
> !(2+2 == 5)
[1] TRUE
```

In other words, since `2+2 == 5` is a `FALSE` statement, it must be the case that `!(2+2 == 5)` is a `TRUE` one. Essentially, what we’ve really done is claim that “not false” is the same thing as “true”. Obviously, this isn’t really quite right in real life. But R lives in a much more black or white world: for R everything is either true or false. No shades of gray are allowed. We can actually see this much more explicitly, like this:

```
> ! FALSE
[1] TRUE
```

Of course, in our $2 + 2 = 5$ example, we didn’t really need to use “not” `!` and “equals to” `==` as two separate operators. We could have just used the “not equals to” operator `!=` like this:

```
> 2+2 != 5
[1] TRUE
```

But there are many situations where you really do need to use the `!` operator. We’ll see some later on.²²

3.8.3 Storing and using logical data

Up to this point, I’ve introduced *numeric data* (in Sections 3.4 and 3.6) and *character data* (in Section 3.7). So you might not be surprised to discover that these `TRUE` and `FALSE` values that R has been

²²A note for those of you who have taken a computer science class: yes, R does have a function for exclusive-or, namely `xor()`. Also worth noting is the fact that R makes the distinction between element-wise operators `&` and `|` and operators that look only at the first element of the vector, namely `&&` and `||`. To see the distinction, compare the behaviour of a command like `c(F,T) & c(T,T)` to the behaviour of something like `c(F,T) && c(T,T)`. If this doesn’t mean anything to you, ignore this footnote entirely. It’s not important for the content of this book.

producing are actually a third kind of data, called *logical data*. That is, when I asked R if `2 + 2 == 5` and it said [1] FALSE in reply, it was actually producing information that we can store in variables. For instance, I could create a variable called `is.the.Party.correct`, which would store R's opinion:

```
> is.the.Party.correct <- 2 + 2 == 5
> is.the.Party.correct
[1] FALSE
```

Alternatively, you can assign the value directly, by typing `TRUE` or `FALSE` in your command. Like this:

```
> is.the.Party.correct <- FALSE
> is.the.Party.correct
[1] FALSE
```

Better yet, because it's kind of tedious to type `TRUE` or `FALSE` over and over again, R provides you with a shortcut: you can use `T` and `F` instead (but it's case sensitive: `t` and `f` won't work).²³ So this works:

```
> is.the.Party.correct <- F
> is.the.Party.correct
[1] FALSE
```

but this doesn't:

```
> is.the.Party.correct <- f
Error: object 'f' not found
```

3.8.4 Vectors of logicals

The next thing to mention is that you can store vectors of logical values in exactly the same way that you can store vectors of numbers (Section 3.6) and vectors of text data (Section 3.7). Again, we can define them directly via the `c()` function, like this:

```
> x <- c(TRUE, TRUE, FALSE)
> x
[1] TRUE TRUE FALSE
```

or you can produce a vector of logicals by applying a logical operator to a vector. This might not make a lot of sense to you, so let's unpack it slowly. First, let's suppose we have a vector of numbers (i.e., a "non-logical vector"). For instance, we could use the `sales.by.month` vector that we were using in Section 3.6. Suppose I wanted R to tell me, for each month of the year, whether I actually sold a book in that month. I can do that by typing this:

```
> sales.by.month > 0
[1] FALSE TRUE TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE
```

and again, I can store this in a vector if I want, as the example below illustrates:

```
> any.sales.this.month <- sales.by.month > 0
> any.sales.this.month
[1] FALSE TRUE TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE
```

²³Warning! `TRUE` and `FALSE` are reserved keywords in R, so you can trust that they always mean what they say they do. Unfortunately, the shortcut versions `T` and `F` do not have this property. It's even possible to create variables that set up the reverse meanings, by typing commands like `T <- FALSE` and `F <- TRUE`. This is kind of insane, and something that is generally thought to be a design flaw in R. Anyway, the long and short of it is that it's safer to use `TRUE` and `FALSE`.

In other words, `any.sales.this.month` is a logical vector whose elements are `TRUE` only if the corresponding element of `sales.by.month` is greater than zero. For instance, since I sold zero books in January, the first element is `FALSE`.

3.8.5 Applying logical operation to text

In a moment (Section 3.9) I'll show you why these logical operations and logical vectors are so handy, but before I do so I want to very briefly point out that you can apply them to text as well as to logical data. It's just that we need to be a bit more careful in understanding how R interprets the different operations. In this section I'll talk about how the equal to operator `==` applies to text, since this is the most important one. Obviously, the not equal to operator `!=` gives the exact opposite answers to `==` so I'm implicitly talking about that one too, but I won't give specific commands showing the use of `!=`. As for the other operators, I'll defer a more detailed discussion of this topic to Section 7.8.5.

Okay, let's see how it works. In one sense, it's very simple. For instance, I can ask R if the word "`cat`" is the same as the word "`dog`", like this:

```
> "cat" == "dog"
[1] FALSE
```

That's pretty obvious, and it's good to know that even R can figure that out. Similarly, R does recognise that a "`cat`" is a "`cat`":

```
> "cat" == "cat"
[1] TRUE
```

Again, that's exactly what we'd expect. However, what you need to keep in mind is that R is not at all tolerant when it comes to grammar and spacing. If two strings differ in any way whatsoever, R will say that they're not equal to each other, as the following examples indicate:

```
> " cat" == "cat"
[1] FALSE
> "cat" == "CAT"
[1] FALSE
> "cat" == "c a t"
[1] FALSE
```

3.9 _____

Indexing vectors

One last thing to add before finishing up this chapter. So far, whenever I've had to get information out of a vector, all I've done is typed something like `months[4]`; and when I do this R prints out the fourth element of the `months` vector. In this section, I'll show you two additional tricks for getting information out of the vector.

3.9.1 Extracting multiple elements

One very useful thing we can do is pull out more than one element at a time. In the previous example, we only used a single number (i.e., `2`) to indicate which element we wanted. Alternatively, we can use a

vector. So, suppose I wanted the data for February, March and April. What I could do is use the vector `c(2,3,4)` to indicate which elements I want R to pull out. That is, I'd type this:

```
> sales.by.month[ c(2,3,4) ]  
[1] 100 200 50
```

Notice that the order matters here. If I asked for the data in the reverse order (i.e., April first, then March, then February) by using the vector `c(4,3,2)`, then R outputs the data in the reverse order:

```
> sales.by.month[ c(4,3,2) ]  
[1] 50 200 100
```

A second thing to be aware of is that R provides you with handy shortcuts for very common situations. For instance, suppose that I wanted to extract everything from the 2nd month through to the 8th month. One way to do this is to do the same thing I did above, and use the vector `c(2,3,4,5,6,7,8)` to indicate the elements that I want. That works just fine

```
> sales.by.month[ c(2,3,4,5,6,7,8) ]  
[1] 100 200 50 0 0 0 0
```

but it's kind of a lot of typing. To help make this easier, R lets you use `2:8` as shorthand for `c(2,3,4,5,6,7,8)`, which makes things a lot simpler. First, let's just check that this is true:

```
> 2:8  
[1] 2 3 4 5 6 7 8
```

Next, let's check that we can use the `2:8` shorthand as a way to pull out the 2nd through 8th elements of `sales.by.months`:

```
> sales.by.month[2:8]  
[1] 100 200 50 0 0 0 0
```

So that's kind of neat.

3.9.2 Logical indexing

At this point, I can introduce an extremely useful tool called **logical indexing**. In the last section, I created a logical vector `any.sales.this.month`, whose elements are `TRUE` for any month in which I sold at least one book, and `FALSE` for all the others. However, that big long list of `TRUEs` and `FALSEs` is a little bit hard to read, so what I'd like to do is to have R select the names of the `months` for which I sold any books. Earlier on, I created a vector `months` that contains the names of each of the months. This is where logical indexing is handy. What I need to do is this:

```
> months[ sales.by.month > 0 ]  
[1] "February" "March"    "April"     "May"
```

To understand what's happening here, it's helpful to notice that `sales.by.month > 0` is the same logical expression that we used to create the `any.sales.this.month` vector in the last section. In fact, I could have just done this:

```
> months[ any.sales.this.month ]  
[1] "February" "March"    "April"     "May"
```

and gotten exactly the same result. In order to figure out which elements of `months` to include in the output, what R does is look to see if the corresponding element in `any.sales.this.month` is `TRUE`. Thus, since element 1 of `any.sales.this.month` is `FALSE`, R does not include "January" as part of the output; but since element 2 of `any.sales.this.month` is `TRUE`, R does include "February" in the output. Note that there's no reason why I can't use the same trick to find the actual sales numbers for those months. The command to do that would just be this:

```
> sales.by.month [ sales.by.month > 0 ]
[1] 100 200 50 25
```

In fact, we can do the same thing with text. Here's an example. Suppose that – to continue the saga of the textbook sales – I later find out that the bookshop only had sufficient stocks for a few months of the year. They tell me that early in the year they had "`high`" stocks, which then dropped to "`low`" levels, and in fact for one month they were "`out`" of copies of the book for a while before they were able to replenish them. Thus I might have a variable called `stock.levels` which looks like this:

```
> stock.levels
[1] "high" "high" "low" "out" "out" "high" "high" "high" "high" "high"
[12] "high"
```

Thus, if I want to know the months for which the bookshop was out of my book, I could apply the logical indexing trick, but with the character vector `stock.levels`, like this:

```
> months[stock.levels == "out"]
[1] "April" "May"
```

Alternatively, if I want to know when the bookshop was either low on copies or out of copies, I could do this:

```
> months[stock.levels == "out" | stock.levels == "low"]
[1] "March" "April" "May"
```

or this

```
> months[stock.levels != "high"]
[1] "March" "April" "May"
```

Either way, I get the answer I want.

At this point, I hope you can see why logical indexing is such a useful thing. It's a very basic, yet very powerful way to manipulate data. We'll talk a lot more about how to manipulate data in Chapter 7, since it's a critical skill for real world research that is often overlooked in introductory research methods classes (or at least, that's been my experience). It does take a bit of practice to become completely comfortable using logical indexing, so it's a good idea to play around with these sorts of commands. Try creating a few different variables of your own, and then ask yourself questions like "how do I get R to spit out all the elements that are [blah]". Practice makes perfect, and it's only by practicing logical indexing that you'll perfect the art of yelling insults at R.²⁴

²⁴Well, I say that... but in my personal experience it wasn't until I started learning "regular expressions" that my loathing of computers reached its peak.

3.10

Quitting R

There's one last thing I should cover in this chapter: how to quit R. When I say this, I'm not trying to apply that R is some kind of pathological addition and that you need to call the R QuitLine or wear patches to control the cravings (although you certainly might argue that there's something seriously pathological about being addicted to R). I just mean how to exit the program. Assuming you're running R in the usual way (i.e., through Rstudio or the default GUI on a Windows or Mac computer), then you can just shut down the application in the normal way. However, R also has a function, called `q()` that you can use to quit. Regardless, when you shut down R it will probably ask you if you want to save the "workspace image". We'll talk a lot more about loading and saving data in Section 4.5, but I figured we'd better quickly cover this now otherwise you're going to get annoyed when you close R at the end of the chapter. Usually you'll see a dialog box come up on your computer asking the question, but here's the text based version that you'd see if you were running R in the old-school way, without any of these fancy graphical options...

```
> q()
Save workspace image? [y/n/c]:
```

The `y/n/c` part here is short for "yes / no / cancel". Type `y` if you want to save, `n` if you don't, and `c` if you've changed your mind and you don't want to quit after all.

What does this actually *mean*? What's going on is that R wants to know if you want to save all those variables that you've been creating, so that you can use them later. This sounds like a great idea, so it's really tempting to hit `y`. However, I very rarely do this... what R is *really* asking is if you want it to store these variables in a "default" data file, which it will automatically reload for you next time you open R. And quite frankly, if I'd wanted to save the variables, then I'd have already saved them before trying to quit. Not only that, I'd have saved them to a location of *my* choice, so that I can find it again later. So I personally never actually hit `y`. In fact, every time I install R on a new machine one of the first things I do is change the settings so that it never asks me again (you can do this in Rstudio really easily: use the menu system to find the "Preferences" option; the dialog box that comes up will give you an option to tell R never to whine about this again).

3.11

Summary

- *Getting started.* We downloaded and installed R and Rstudio (Section 3.1).
- *Basic commands.* We talked a bit about the logic of how works and in particular how to type commands into the R console (Section 3.2), and in doing so learned how to perform basic calculations using the arithmetic operators `+`, `-`, `*`, `/` and `^`. (Section 3.3)
- *Introduction to functions.* We saw several different functions, three that are used to perform numeric calculations (`sqrt()`, `abs()`, `round()`; Section 3.5), one that applies to text (`nchar()`; Section 3.7.1), and one that works on any variable (`length()`; Section 3.6.5). In doing so, we talked a bit about how argument names work, and learned about default values for arguments. (Section 3.5.1)
- *Introduction to variables.* We learned the basic idea behind variables, and how to assign values to variables using the assignment operator `<-` (Section 3.4). We also learned how to create vectors using the combine function `c()`. (Section 3.6)

- *Data types.* Learned the distinction between numeric, character and logical data; including the basics of how to enter and use each of them. (Sections 3.4 to 3.8)
- *Logical operations.* Learned how to use the logical operators `==`, `!=`, `<`, `>`, `<=`, `=>`, `!`, `&` and `|`. (Section 3.8). And learned how to use logical indexing. (Section 3.9)

4. Additional R concepts

In Chapter 3 our main goal was to get started in R. We covered a lot of the basics and started getting a feel for how R works. As we go through the book we'll run into a lot of new R concepts, which I'll explain alongside the relevant data analysis concepts. However, there's still quite a few things that I need to talk about now, otherwise we'll run into problems when we start trying to work with data and do statistics. So that's the goal in this chapter: to build on the introductory content from the last chapter, to get you to the point that we can start using R for statistics. Broadly speaking, the chapter comes in two parts. The first half of the chapter is devoted to the “mechanics” of R: installing and loading packages, managing the workspace, navigating the file system, and loading and saving data. In the second half, I'll talk more about what kinds of variables exist in R, and introduce three new kinds of variables: factors, data frames and formulas. I'll finish up by talking a little bit about the help documentation in R as well as some other avenues for finding assistance. In general, I'm not trying to be comprehensive in this chapter, I'm trying to make sure that you've got the basic foundations needed to tackle the content that comes later in the book. However, a lot of the topics are revisited in more detail later, especially in Chapters 7 and 8.

4.1

Using comments

Before discussing any of the more complicated stuff, I want to introduce the `comment` character, `#`. It has a simple meaning: it tells R to ignore everything else you've written on this line. You won't have much need of the `#` character immediately, but it's very useful later on when writing scripts (see Chapter 8). However, while you don't need to use it, I want to be able to include comments in my R extracts. For instance, if you read this:¹

```
> seeker <- 3.1415      # create the first variable
> lover <- 2.7183       # create the second variable
> keeper <- seeker * lover # now multiply them to create a third one
> print( keeper )        # print out the value of 'keeper'
[1] 8.539539
```

it's a lot easier to understand what I'm doing than if I just write this:

```
> seeker <- 3.1415
> lover <- 2.7183
> keeper <- seeker * lover
```

¹Notice that I used `print(keeper)` rather than just typing `keeper`. From now on I'll start to use the `print()` function to display things because I think it helps make clear what I'm doing, but in practice people rarely do this.

```
> print( keeper )
[1] 8.539539
```

So, from now on, you'll start seeing some `#` characters appearing in the extracts, with some human-readable explanatory remarks next to them. These are still perfectly legitimate commands, since R knows that it should ignore the `#` character and everything after it. But hopefully they'll help make things a little easier to understand.

4.2

Installing and loading packages

In this section I discuss R **packages**, since almost all of the functions you might want to use in R come in packages. A package is basically just a big collection of functions and other R objects that are all grouped together under a common name. Some packages are already installed when you put R on your computer, but the vast majority of them of R packages are out there on the internet, waiting for you to download, install and use them.

4.2.1 What packages do you have installed?

When you download R initially, there are about 30 or so packages that are already installed. You can find out what packages are installed on your computer by typing `.packages(TRUE)` at the command line. You should see something very similar to this:

```
> .packages(TRUE)
[1] "base"      "boot"       "class"      "cluster"
[5] "codetools"  "compiler"   "datasets"   "foreign"
[9] "graphics"   "grDevices"  "grid"       "KernSmooth"
[13] "lattice"    "MASS"       "Matrix"     "methods"
[17] "mgcv"       "nlme"       "nnet"       "rpart"
[21] "spatial"    "splines"    "stats"     "stats4"
[25] "survival"   "tcltk"     "tools"     "utils"
[29] "manipulate"
```

Alternatively, if you want a bit more detail about what each package does, you can try typing

```
> library()
```

This command produces a much longer output that gives the name of every installed package, as well as a short description of what the package does. Not only that, it tells you whereabouts on your computer all the files are stored, which can be useful. If you're using Rstudio this will show up in the “source” panel rather than getting printed out in the console. This output goes on at some length, and at this early stage none of it really means very much to you, so I haven't bothered to include it here.

4.2.2 What packages do you have loaded?

So there are about 30 or so packages that you already have on your computer. However, if you want to make use of some of the functions or data in a particular package, it's not sufficient just to have the

package on your computer. The package must be *loaded*. When you start up R it will load some packages, but not all of them. To see which packages are loaded, we can use the `search()` function.

```
> search()
[1] ".GlobalEnv"      "tools:rstudio"    "package:stats"
[4] "package:graphics" "package:grDevices" "package:utils"
[7] "package:datasets" "package:methods"   "Autoloads"
[10] "package:base"
```

This output is showing you something called the **search path**, and it's slightly more than just a list of packages, but we'll talk more about it later (Section 7.12.3). For now, though, let's just focus on the packages: what this is telling me is that, right now, I have access to the `stats`, `graphics`, `grDevices`, `utils`, `datasets`, `methods` and `base` packages. Any function, variable or data set that is stored in one of these packages is accessible to R.

We can illustrate this point using the `find()` function. Suppose that there's some R function or variable whose name you know,² and you're pretty sure that it's in one of the packages that you've got loaded, but you don't know which one. The `find()` function can search through the packages and find it for you. The `find` function has a number of arguments, but the only one we'll use is `what`: a character string giving the name of the function that you're looking for. To give a concrete example, one of the functions that we'll need to use a lot (especially in Chapter 13) is `t.test()`. Let's find it:

```
> find( what = "t.test" )
[1] "package:stats"
```

Okay, so R is clearly telling me that it thinks that the `t.test()` function exists, and that it's stored in the `stats` package.³ Alternatively, suppose I were to ask R to find something called `cancer`. Let's see what happens:

```
> find( what = "cancer" )
character(0)
```

This doesn't look like the name of a package, does it? What you're seeing here is R telling you that the output of the command is an empty string: that is, it's returned a value that is notionally a kind of text (i.e., “character” data), but it doesn't actually have any text in it! Every now and then you'll see R do this, so I wanted to highlight this behaviour so that you know what it means. But in terms of our original goal, which was to try to find something called `cancer`, what this response means is that R looked through all those packages and didn't find it.

4.2.3 How do you load a package?

The previous example highlights an important point. Just because the thing you're looking for doesn't exist in any of the loaded packages doesn't mean that it doesn't exist somewhere. In fact, there is something in R called `cancer`. It's a data set stored in the `survival` package. And if you compare the output of the `.packages(TRUE)` command that we used in Section 4.2.1) to the output of the `search()`

²If you don't know the exact name of the function, you can search for functions that partially match against your input. The function that you use to do this is called `apropos()`, but we won't touch it in this book because it relies on a very powerful (but complicated) text processing tool called *regular expressions*. Regular expressions totally rock (sort of: they get on my nerves some days), but you need to work your way up to them! Because this is tricky, later on (in Section 4.12) I'm going to suggest an easier solution: use Google, or www.rseek.org to do the searching for you.

³Note that I typed "`t.test`" not "`t.test()`". This is because the *proper* name for the function is `t.test`. Throughout this book I'm going to add the parentheses because I think it's a useful visual convention, but it's worth remembering that the actual *function name* itself doesn't include them. The parentheses are part of the *command* that you type.

command that we typed in Section 4.2.2), you'll notice that I actually *do* have the `survival` package: it's installed on my computer, but R hasn't loaded it. This leads to the natural question... how do we load a package?

Loading packages is pretty easy. The function that we need to use to load a package is called `library()`. The logic behind the name is that R stores all your packages in a "library" of packages, so you can kind of imagine that you're going to the library and picking out a package.⁴ In any case, the only argument that we need to care about for our purposes is `package` which you use to specify the name of the package you want to load. No surprises there, so here's what I type:

```
> library( package = "survival" )
```

When you type a command like this, sometimes you'll see output and sometimes you won't. Most likely, if you're reading through this book for the first time and you haven't been customising R at all, you will see this as the output:

```
Loading required package: splines
```

What this means is that R has actually loaded *two* packages. It loaded `survival`, because that's the one you asked for, but it's also loaded the `splines` package, because it turns out that the `survival` package relies on some of the tools provided by `splines`. A little later we'll have a look at some of the things that R tends to print out when you load packages. In the meantime, if we use the `search()` function to have a look at what packages R we've got loaded, here's what we get:

```
> search()
[1] ".GlobalEnv"      "package:survival"   "package:splines"
[4] "tools:rstudio"    "package:stats"     "package:graphics"
[7] "package:grDevices" "package:utils"     "package:datasets"
[10] "package:methods"   "Autoloads"       "package:base"
```

Notice that we've added two new entries here.

One very useful thing to remember about the `library()` function, *especially* since you'll use it a lot, is that `package` is the first argument, so you don't actually have to bother specifying it by name. Even better, the function is written in a pretty clever way, and as a result you don't need to include the quote marks (that's *not* true in general though!). So in practice, the way you'd load the package would be to type this:

```
> library( survival )
```

In any case, now that we've loaded the `survival` package, R now acknowledges the existence⁵ of `cancer`. To see this, let's use the `find()` function again. This time I won't bother naming my arguments, because as it happens the `what` argument is the first one:

```
> find( "cancer" )
[1] "package:survival"
```

4.2.4 How do you unload a package?

Sometimes, especially after a long session of working with R, you find yourself wanting to get rid of some of those packages that you've loaded. I'll show you two ways to do it. Firstly, there's the `detach()`

⁴Yeah, I know, it's a pretty clunky metaphor. In fact, I've heard tell that future versions of R will introduce a new function called `use()` that you can use instead of `library()`, which will then be slowly phased out.

⁵See also the `exists()` function.

function. This is the main way to do it, but it's slightly counterintuitive since it doesn't use the same arguments as the `library()` function. The main argument we care about is the `name` of the package to be detached, but what you need to input is a character string that matches one of the things we saw when we called the `search()` function. That is, if we want to unload the `survival` package, what we need to do is this:

```
> detach( name = "package:survival" )
```

And although it's possible to drop the `name =` part, since `name` is the first argument to the `detach()` function, you really do need to specify the full `"package:survival"` string. So, to unload the `splines` package, I could do this:

```
> detach("package:splines")
```

but I wouldn't be able to shorten that command any further. There is a good reason for this: there are situations where you have to `detach()` things that aren't packages, and so the `detach()` function needs to have this extra detail. But it still bugs me a little. To that end, I've written my own function called `unlibrary()`. It's in the `lsr` package so you can't use it until you've downloaded, installed and loaded `lsr`, but I figured that I should mention it here. The `unlibrary()` function lets you unload a package using the same kind of command that you used to load it in the first place. So, if I'd had the `lsr` package⁶ loaded, what I could have done instead is this:

```
> unlibrary( survival )
```

4.2.5 A few extra comments

Sections 4.2.3 and 4.2.4 cover the main things you need to know about loading and unloading packages. However, there's a couple of other details that I want to draw your attention to. A concrete example is the best way to illustrate. One of the other packages that you already have installed on your computer is the `Matrix` package, so let's load that one and see what happens:

```
> library( Matrix )
Loading required package: lattice

Attaching package: 'Matrix'

The following object(s) are masked from 'package:base':

det
```

This is slightly more complex than the output that we got last time, but it's not too complicated. In the first line of the output, we see that in addition to loading the `Matrix` package, R has also loaded the `lattice` package. So far that's the same sort of thing that we saw last time. But there's something new here. R tells us that it has "masked" something from the `base` package, something called "det". In case you care, `det()` is a function used to calculate a matrix determinant (a useful thing in advanced contexts, but of no real interest to us here), but that's besides the point. What I really want to focus on is this "masking" business. To understand what's happening here, let's use the `find()` function to see where the `det()` function is. When we do this...

```
> find( "det" )
[1] "package:Matrix" "package:base"
```

⁶The name comes from the name of the book: "Learning Statistics with R". The fact that it happens to be a homophone for "loser" is a complete co-incidence. Why do you ask?

... we discover that there's actually *two* `det()` functions, one in the `base` package and another in the `Matrix` package. What's happened here is that, when we loaded the `Matrix` package, R noticed that there are now two functions with the same name, and it's printed out a warning. Specifically, what the "masking" bit means is that if you try to use the `det()` function, R will use the one from the `Matrix` package, not the one from the `base` package.⁷

Okay, one more thing to point out. Recall that the `Matrix` package requires the `lattice` package, so when I loaded `Matrix` R also loaded `lattice`. What would happen if I tried to unload the `lattice` package, without first having unloaded `Matrix`? Let's see:

```
> detach( "package:lattice" )
Error: package lattice is required by Matrix so will not be detached
```

R refuses to do it. This can be quite useful, since it stops you from accidentally removing something that you still need. So, if I want to remove both `Matrix` and `lattice`, I need to do it in the correct order, like this:

```
> detach( "package:Matrix" )
> detach( "package:lattice" )
```

4.2.6 Downloading new packages

One of the main selling points for R is that there are thousands of packages that have been written for it, and these are all available online. So whereabouts online are these packages to be found, and how do we download and install them? There is a big repository of packages called the "Comprehensive R Archive Network" (CRAN), and the easiest way of getting and installing a new package is from one of the many CRAN mirror sites. Conveniently for us, R provides a function called `install.packages()` that you can use to do this. However, before we use it, it's a good idea to set your preferred mirror, so that R knows which of the various CRAN sites it should try to download from. The easiest way to do this is to use the Rstudio preferences. On a Windows computer, select the Options entry in the Tools menu. On a Mac, select the Preferences option in the Rstudio menu. This opens up an Options window (shown in Figure 4.1). Among the "R General" options there's a spot where you can choose your CRAN mirror. Click the "Change" button, and it will bring up a list of CRAN mirrors. Select the one you want, and then hit okay. You've now selected your preferred CRAN mirror, and now that you've done so, R won't bug you about what mirror to use.

Assuming that you've set your preference, here's how we would use the `install.packages()` function. There are a few arguments that we might want to specify:

- `pkgs`. This is a string that contains the name of the package, or (if you want to install more than one package at once) a vector of strings containing the names of all the packages you want to install.
- `dependencies`. This is a logical value indicating whether or not R should *also* install any packages that are required by those listed in `pkgs`. The default value is `FALSE` (with good reason), but I usually set it to `TRUE` to make sure that I get all the packages I need.
- `repos`. This is a string that contains the website of your CRAN mirror. By default, R checks the your "options" to see what it should use. In other words, the default value of `repos` is identical to whatever the output of the command `getOption("repos")` happens to be. As long as you've set your CRAN mirror already, you don't need to bother with this one at all.

⁷Tip for advanced users. You can get R to use the one from the `base` package by using `base:::det()` as your command rather than `det()`, since the `base:::` part tells R explicitly which package to use. See also `:::`: if you're especially keen to force R to do things it otherwise wouldn't, but take care, since `:::` can be dangerous.

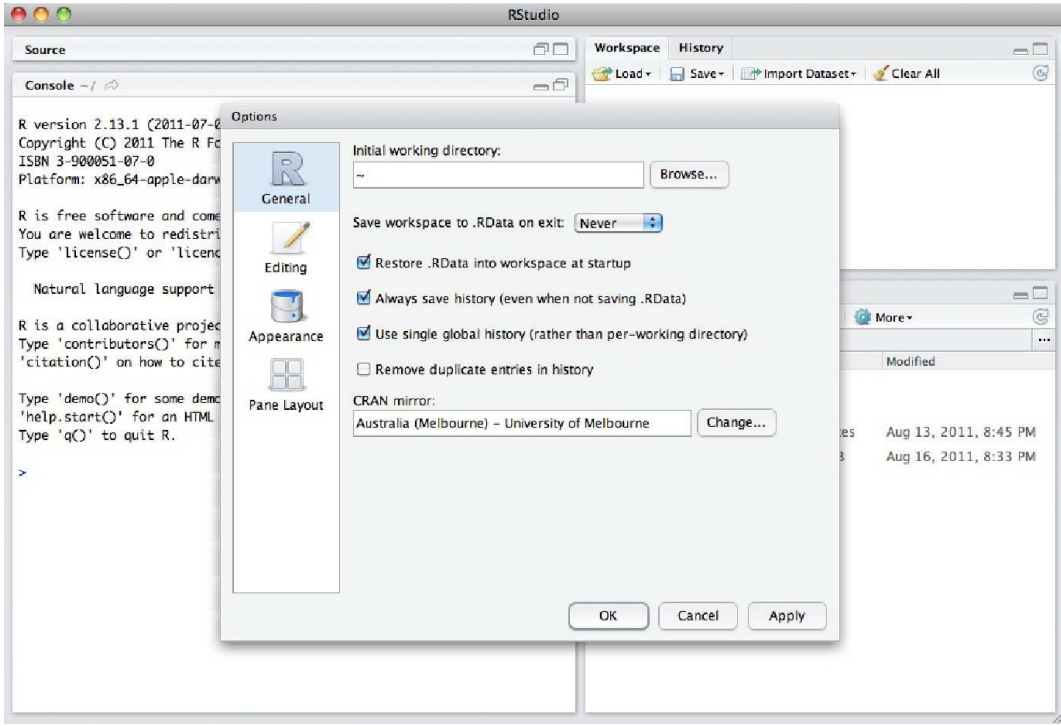


Figure 4.1: The options dialog box. To change the CRAN mirror, you just click the “Change...” button and then select a mirror from the list. Note that there are quite a few other settings in there that you might be interested in changing.

- **lib**. This is a string that specifies the location on your computer where the package should be installed into. Again, R has a pretty sensible default here, so you don’t need to bother with this argument at all until you’re feeling a bit more comfortable with R.

One of the packages that we’ll use in this book is the `car` package (short for “Companion to Applied Regression”, which is the title of an excellent book that I’d strongly recommend reading once you’re done with this one). To install it, I’ll use the following command:

```
> install.packages( pkgs = "car", dependencies = TRUE )
```

The output that we get from this is fairly complicated, so I’ll show it to you line by line and explain what each part means. Firstly, we get something that looks like this:

```
Installing package(s) into /Library/Frameworks/R.framework/Versions/2.15/Resources/library
(as lib is unspecified)
```

What this is telling you is that R has noticed that I didn’t specify the `lib` argument, so it’s decided to store the package in the same place that all the rest of my R packages are stored. Next comes this:

```
trying URL 'http://cran.csiro.au/bin/macosx/leopard/contrib/2.15/car_2.0-15.tgz'
Content type 'application/x-gzip' length 1064620 bytes (1.0 Mb)
opened URL
```

```
=====
downloaded 1.0 Mb
```

This is a whole lot of information in which R first tells you exactly whereabouts on the internet it's going to try to download from. The second line is a standard piece of "header" information that all internet communications provide: that is, before you download something from a website, the website provides your computer with a little bit of data about what type of data it's downloading, and how much of it there is. In the third line, R reports that it has successfully started downloading; and it then displays all those = symbols as a kind of text-based progress bar, to indicate that it is actually receiving the data; and then finally it tells you that it has succeeded in downloading all the data. Finally, we get something like this:

```
The downloaded binary packages are in  
/var/folders/zt/stn3df0n7tndrdkyxzdwp8m0000gn/T//RtmpJVGQv9/downloaded_packages
```

which is just some information about where it saved all the temporary files. At this point, the package has been installed. If it had turned our that I needed to install a lot of dependencies, then this sequence would be repeated several times automatically: it's ugly to watch, but thankfully you don't have to do anything other than watch. Actually, you don't have to even do that if you close your eyes.

4.2.7 Updating R and R packages

Every now and then the authors of packages release updated versions. The updated versions often add new functionality, fix bugs, and so on. It's generally a good idea to update your packages periodically. The easiest way to do that is to type

```
> update.packages()
```

at the command line. When you enter this command, R will look online to see which of your packages have newer versions available. It won't automatically upgrade the packages, but will ask you to make a decision for each package that it thinks you can upgrade (this can take a while if you've got a lot of packages installed and haven't updated in a while). Once you've indicated which ones you want updated, R will then go through the same process that it went through when installing the packages in the first place.

About every six months or so, a new version of R is released. To get the new version you can get to the CRAN website and download the most recent version of R, and install it in the same way you did when you originally installed R on your computer. This used to be a slightly frustrating event, because whenever you downloaded the new version of R, you would lose all the packages that you'd downloaded and installed, and would have to repeat the process of re-installing them. This was pretty annoying, and there were some neat tricks you could use to get around this. However, it looks like newer versions of R no longer suffer from this problem. However, just in case you ever need to use it, here's the trick I used. First, before you update R, use the `.packages(TRUE)` command to get a list of all your installed packages, and save it to a file:

```
> packages <- .packages(TRUE)  
> save(packages,file="mypackages.Rdata")
```

Then, once you've installed the new version of R, load the list of packages, and use the `install.packages()` function to fetch them all from CRAN:

```
> load("mypackages.Rdata")  
> install.packages(packages,dependencies=TRUE)
```

This method does still mean that you need to download all the packages again, but the important thing is that you don't have to manually type in all the package names.

4.2.8 What packages does this book use?

There are several packages that I make use of in this book. The most prominent of them are as follows:

- **lsr**. This is the *Learning Statistics with R* package that accompanies this book. It doesn't have a lot of interesting high-powered tools: it's just a small collection of handy little things that I think can be useful to novice users. As you get more comfortable with R this package should start to feel pretty useless to you.
- **psych**. This package, written by William Revelle, includes a lot of tools that are of particular use to psychologists. In particular, there's several functions that are particularly convenient for producing analyses or summaries that are very common in psych, but less common in other disciplines.
- **car**. This is the *Companion to Applied Regression* package, which accompanies the excellent book of the same name by [Fox and Weisberg \(2011\)](#). It provides a lot of very powerful tools, only some of which we'll touch in this book.

Besides these three, there are a number of packages that I use in a more limited fashion: **gplots**, **sciplot**, **foreign**, **effects**, **TeachingDemos**, **R.matlab**, **gdata**, **lntest**, and probably one or two others that I've missed. There are also a number of packages that I refer to but don't actually use in this book, such as **reshape**, **compute.es**, **HistData** and **multcomp** among others. Finally, there are a number of packages that provide more advanced tools that I hope to talk about in future versions of the book, such as **sem**, **ez**, **nlme** and **lme4**. In any case, whenever I'm using a function that isn't in the core packages, I'll make sure to note this in the text.

4.3 _____

Managing the workspace

Let's suppose that you're reading through this book, and what you're doing is sitting down with it once a week and working through a whole chapter in each sitting. Not only that, you've been following my advice and typing in all these commands into R. So far during this chapter, you'd have typed quite a few commands, although the only ones that actually involved creating variables were the ones you typed during Section 4.1. As a result, you currently have three variables; **seeker**, **lover**, and **keeper**. These three variables are the contents of your **workspace**. The workspace is a key concept in R, so in this section we'll talk a lot about what it is and how to manage its contents.

4.3.1 Listing the contents of the workspace

The first thing that you need to know how to do is list the contents of the workspace. Although it's pretty obvious what our variables are right now, it becomes harder to remember when you're doing real data analysis. To do this you can use the **objects()** function.

```
> objects()
[1] "keeper" "lover"  "seeker"
```

Of course, in the true R tradition, the `objects()` function has a lot of fancy capabilities that I'm glossing over in this example, and moreover there are also several other functions that you can use, including `ls()` which is pretty much identical to `objects()`, and `ls.str()` which you can use to get a fairly detailed description of all the variables in the workspace. In fact, the `lsr` package actually includes its own function that you can use for this purpose, called `who()`. The reason for using the `who()` function is pretty straightforward: in my everyday work I find that the output produced by the `objects()` command isn't *quite* informative enough, because the only thing it prints out is the name of each variable; but the `ls.str()` function is *too* informative, because it prints out a lot of additional information that I really don't like to look at. The `who()` function is a compromise between the two. First, now that we've got the `lsr` package installed, we need to load it:

```
> library(lsrr)
```

and now we can use the `who()` function:

```
> who()
-- Name --  -- Class --  -- Size --
keeper      numeric      1
lover       numeric      1
seeker      numeric      1
```

As you can see, the `who()` function lists all the variables and provides some basic information about what kind of variable each one is and how many elements it contains. Personally, I find this output much easier more useful than the very compact output of the `objects()` function, but less overwhelming than the extremely verbose `ls.str()` function.

4.3.2 Removing variables from the workspace

Looking over that list of variables, it occurs to me that I really don't need them any more. I created them originally just to make a point, but they don't serve any useful purpose anymore, and now I want to get rid of them. I'll show you how to do this, but first I want to warn you – there's no “undo” option for variable removal. Once a variable is removed, it's gone forever unless you save it to disk. I'll show you how to do *that* in Section 4.5, but quite clearly we have no need for these variables at all, so we can safely get rid of them.

This is where the `remove` function `rm()` comes in handy. The simplest way to use `rm()` is just to type in a (comma separated) list of all the variables you want to remove. Let's say I want to get rid of `seeker` and `lover`, but I would like to keep `keeper`. To do this, all I have to do is type:

```
> rm( seeker, lover )
```

There's no visible output, but if I now inspect the workspace

```
> who()
-- Name --  -- Class --  -- Size --
keeper      numeric      1
```

I see that there's only the `keeper` variable left. As you can see, `rm()` can be very handy for keeping the workspace tidy.

A slightly different thing you often find yourself wanting to do is to wipe the slate clean: to remove *all* of the variables in the workspace. Obviously, at the moment I only have the one variable, but let's suppose I've just typed `junk1 <- "blah"`, `junk2 <- "blah"` and `junk3 <- "blah"`, so now I have four

variable in my workspace. How do I get rid of these? One possibility is to do the same thing as before, typing in the name of every variable, but clearly that's inefficient. A second possibility is provided by the `rm()` function, which has an optional argument called `list`, in which you can input a vector of strings that contains the names of all the variables that you want to delete. Why does that help? Because what we can do is this:

```
> rm( list = objects() )
```

That does work, but I still find that that's way too much typing (yes, I'm lazy). To redress this, the `lsr` package contains a function called `rmAll()` which will delete all variables in the workspace, though it does ask you for confirmation before deleting everything...

```
> rmAll()
Current contents of workspace:
[1] "junk1" "junk2" "junk3"
Remove all objects? [y/n] y
```

If we now inspect the workspace, we see that the workspace is empty:

```
> who()
No variables found
```

One final option I should point out if you're using Rstudio, one easy way to remove all of your variables is provided by the “workspace” panel, which by default is one of the tabs located in the top right hand corner of the screen. Notice that it has a button that says “Clear All”? If you click on this, it will remove all variables, in pretty much the same way that `rmAll()` does. Just like `rmAll()` it will ask you to confirm before wiping the workspace clean.

4.4

Navigating the file system

In this section I talk a little about how R interacts with the file system on your computer. It's not a terribly interesting topic, but it's useful. As background to this discussion, I'll talk a bit about how file system locations work in Section 4.4.1. I suspect that this is very familiar to most readers (so you can skip straight to Section 4.4.2), but it probably won't be well understood by everyone, especially since modern operating systems seem to be going out of their way to hide this from you, and not everyone grew up being forced to learn how to navigate around a computer using a DOS or UNIX shell.

4.4.1 The file system itself

In this section I describe the basic idea behind file locations and file paths. Regardless of whether you're using Window, Mac OS or Linux, every file on the computer is assigned a (fairly) human readable address, and every address has the same basic structure: it describes a *path* that starts from a *root* location, through a series of *folders* (or if you're an old-school computer user, *directories*), and finally ends up at the file.

On a Windows computer the root is the physical drive⁸ on which the file is stored, and for most home computers the name of the hard drive that stores all your files is C: and therefore most file names on

⁸Well, the partition, technically.

Windows begin with `C:`. After that comes the folders, and on Windows the folder names are separated by a `\` symbol. So, the complete path to this book on my Windows computer might be something like this:

`C:\Users\dan\Rbook\LSR.pdf`

and what that *means* is that the book is called `LSR.pdf`, and it's in a folder called `Rbook` which itself is in a folder called `dan` which itself is ... well, you get the idea. On Linux, Unix and Mac OS systems, the addresses look a little different, but they're more or less identical in spirit. Instead of using the backslash, folders are separated using a forward slash, and unlike Windows, they don't treat the physical drive as being the root of the file system. So, the path to this book on my Mac might be something like this:

`/Users/dan/Rbook/LSR.pdf`

So that's what we mean by the "path" to a file. The next concept to grasp is the idea of a **working directory** and how to change it. For those of you who have used command line interfaces previously, this should be obvious already. But if not, here's what I mean. The working directory is just "whatever folder I'm currently looking at". Suppose that I'm currently looking for files in Explorer (if you're using Windows) or using Finder (on a Mac). The folder I currently have open is my user directory (i.e., `C:\Users\dan` or `/Users/dan`). That's my current working directory.

The fact that we can imagine that the program is "in" a particular directory means that we can talk about moving *from* our current location *to* a new one. What that means is that we might want to specify a new location in relation to our current location. To do so, we need to introduce two new conventions. Regardless of what operating system you're using, we use `.` to refer to the current working directory, and `..` to refer to the directory above it. This allows us to specify a path to a new location in relation to our current location, as the following examples illustrate. Let's assume that I'm using my Windows computer, and my working directory is `C:\Users\dan\Rbook`. The table below shows several addresses in relation to my current one:

absolute path (i.e., from root)	relative path (i.e. from <code>C:\Users\dan\Rbook</code>)
<code>C:\Users\dan</code>	<code>..</code>
<code>C:\Users</code>	<code>..\..</code>
<code>C:\Users\dan\Rbook\source</code>	<code>.\source</code>
<code>C:\Users\dan\nerdstuff</code>	<code>..\nerdstuff</code>

There's one last thing I want to call attention to: the `~` directory. I normally wouldn't bother, but R makes reference to this concept sometimes. It's quite common on computers that have multiple users to define `~` to be the user's home directory. On my Mac, for instance, the home directory `~` for the "dan" user is `\Users\dan\`. And so, not surprisingly, it is possible to define other directories in terms of their relationship to the home directory. For example, an alternative way to describe the location of the `LSR.pdf` file on my Mac would be

`~\Rbook\LSR.pdf`

That's about all you really need to know about file paths. And since this section already feels too long, it's time to look at how to navigate the file system in R.

4.4.2 Navigating the file system using the R console

When you want to load or save a file in R it's important to know what the working directory is. You

can find out by using the `getwd()` command. For the moment, let's assume that I'm using Mac OS or Linux, since there's some subtleties to Windows. Here's what happens:

```
> getwd()
[1] "/Users/dan"
```

We can change the working directory quite easily using `setwd()`. The `setwd()` function has only the one argument, `dir`, is a character string specifying a path to a directory, or a path relative to the working directory. Since I'm currently located at `/Users/dan`, the following two are equivalent:

```
> setwd("/Users/dan/Rbook/data")
> setwd("./Rbook/data")
```

Now that we're here, we can type `list.files()` command to get a listing of all the files in that directory. Since this is the directory in which I store all of the data files that we'll use in this book, here's what we get as the result:

```
> list.files()
[1] "afl24.Rdata"           "agpp.Rdata"          "anscombesquartet.Rdata"
[4] "awesome.Rdata"         "awesome2.Rdata"       "booksales.Rdata"
[7] "booksales.csv"         "booksales2.csv"        "cards.Rdata"
[10] "chapek9.Rdata"        "chico.Rdata"         "clinicaltrial.Rdata"
[13] "drawttesthyp.R"        "drugs.wmc.rt.Rdata"  "happy.Rdata"
[16] "harpo.Rdata"          "harpo2.Rdata"         "parenthood.Rdata"
[19] "parenthood2.Rdata"    "rankcorr.Rdata"       "salem.Rdata"
[22] "simdata.csv"          "zeppo.Rdata"
```

Not terribly exciting, I'll admit, but it's useful to know about. In any case, there's only one more thing I want to make a note of, which is that R also makes use of the home directory. You can find out what it is by using the `path.expand()` function, like this:

```
> path.expand("~")
[1] "/Users/dan"
```

You can change the user directory if you want, but we're not going to make use of it very much so there's no reason to. The only reason I'm even bothering to mention it at all is that when you use Rstudio to open a file, you'll see output on screen that defines the path to the file relative to the `~` directory. I'd prefer you not to be confused when you see it.⁹

4.4.3 Why do the Windows paths use the wrong slash?

Let's suppose I'm on Windows. As before, I can find out what my current working directory is like this:

```
> getwd()
[1] "C:/Users/dan/"
```

This seems about right, but you might be wondering why R is displaying a Windows path using the wrong type of slash. The answer is slightly complicated, and has to do with the fact that R treats the \

⁹One additional thing worth calling your attention to is the `file.choose()` function. Suppose you want to load a file and you don't quite remember where it is, but would like to browse for it. Typing `file.choose()` at the command line will open a window in which you can browse to find the file; when you click on the file you want, R will print out the full path to that file. This is kind of handy.

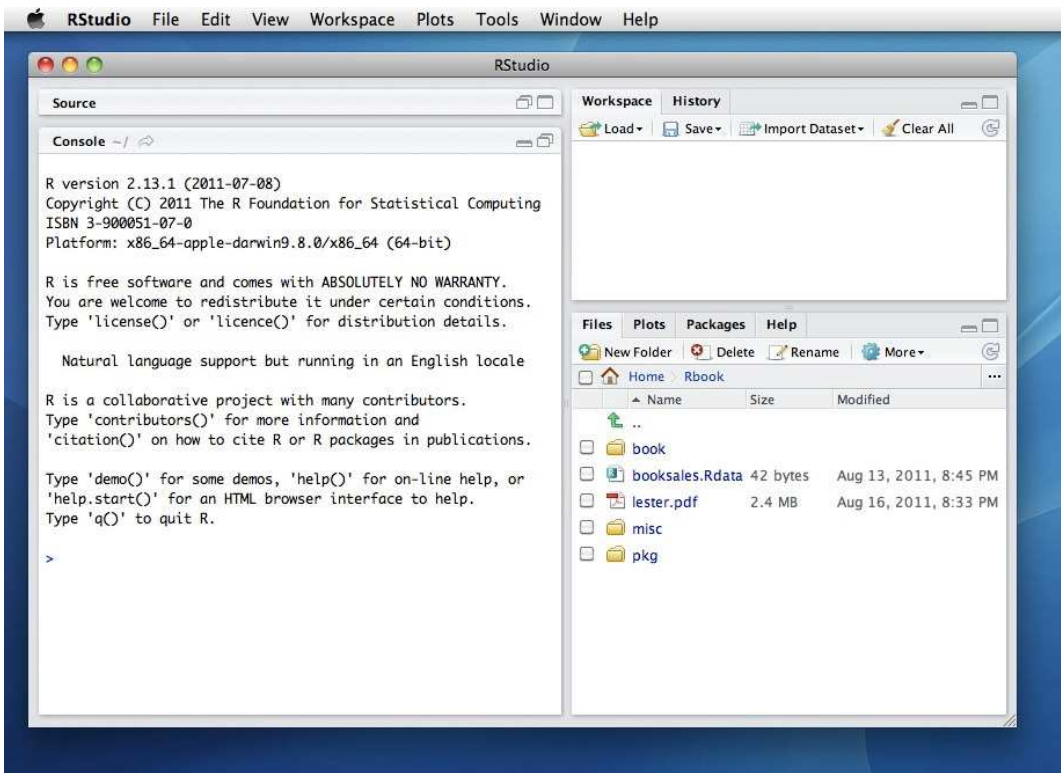


Figure 4.2: The “file panel” is the area shown in the lower right hand corner. It provides a very easy way to browse and navigate your computer using R. See main text for details.

.....

character as “special” (see Section 7.8.7). If you’re deeply wedded to the idea of specifying a path using the Windows style slashes, then what you need to do is to type `\` whenever you mean `\`. In other words, if you want to specify the working directory on a Windows computer, you need to use one of the following commands:

```
> setwd( "C:/Users/dan" )
> setwd( "C:\\Users\\dan" )
```

It’s kind of annoying to have to do it this way, but as you’ll see later on in Section 7.8.7 it’s a necessary evil. Fortunately, as we’ll see in the next section, Rstudio provides a much simpler way of changing directories...

4.4.4 Navigating the file system using the Rstudio “file” panel

Although it’s important to understand how all this command line stuff works, in many (maybe even most) situations there’s an easier way. For our purposes, the easiest way to navigate the file system is to make use of Rstudio’s built in tools. The “file” panel – the lower right hand area in Figure 4.2 – is actually a pretty decent file browser. Not only can you just point and click on the names to move around

the file system, you can also use it to set the working directory, and even load files.

Here's what you need to do to change the working directory using the file panel. Let's say I'm looking at the actual screen shown in Figure 4.2. The file panel is currently showing me the `~\Rbook\` directory (that's what the "Home > Rbook" part is telling me). If I click on the word "book" just below, the file panel will now move to the directory `~\Rbook\`. At the top it will now show "Home > Rbook > book", and the panel itself will now list the contents of the new directory.¹⁰ Importantly, you need to understand that this is *Rstudio* doing the browsing. R itself hasn't done anything. You can tell because there's nothing printed in the console. This is nice, because it means that you can look for files (and, as we'll see in the next section, even load files) without having to change the R working directory. However, sometimes we really do want to change the working directory. To do this, click on the "More" button. This will bring up a little menu, and one of the options will be "Set as Working Directory". If you select that option, then R really will change the working directory. You can tell that it has done so because the console now has this in it:

```
> setwd("~/Rbook/book")
```

In other words, *Rstudio* sends a command to the R console, exactly as if you'd typed it yourself. If you want to move the file panel back up, click on the "..." link in the panel.

As you can tell, the file panel is a very handy little tool for navigating the file system. But it can do more than just navigate. As we'll see later, it can be used to open files. And if you look at the buttons and menu options that it presents, you can even use it to rename, delete, copy or move files, and create new folders. However, since most of that functionality isn't critical to the basic goals of this book, I'll let you discover those on your own.

4.5

Loading and saving data

There are several different types of files that are likely to be relevant to us when doing data analysis. There are three in particular that are especially important from the perspective of this book:

- *Workspace files* are those with a `.Rdata` file extension. This is the standard kind of file that R uses to store data and variables. They're called "workspace files" because you can use them to save your whole workspace.
- *Comma separated value (CSV) files* are those with a `.csv` file extension. These are just regular old text files, and they can be opened with almost any software. It's quite typical for people to store data in CSV files, precisely because they're so simple.
- *Script files* are those with a `.R` file extension. These aren't data files at all; rather, they're used to save a collection of commands that you want R to execute later. They're just text files, but we won't make use of them until Chapter 8.

There are also several other types of file that R makes use of,¹¹ but they're not really all that central to our interests. There are also several other kinds of data file that you might want to import into R. For instance, you might want to open Microsoft Excel spreadsheets (`.xls` files), or data files that have been saved in the native file formats for other statistics software, such as SPSS, SAS, Minitab, Stata or Systat. Finally, you might have to handle databases. R tries hard to play nicely with other software, so

¹⁰By the way, you can also click the "..." button to achieve the same effect: it will bring up a little dialog box that will let you browse to find the folder you want.

¹¹Notably those with `.rda`, `.Rd`, `.Rhistory`, `.rdb` and `.rdx` extensions

it has tools that let you open and work with any of these and many others. I'll discuss some of these other possibilities elsewhere in this book (Section 7.9), but for now I want to focus primarily on the two kinds of data file that you're most likely to need: `.Rdata` files and `.csv` files.

4.5.1 Loading workspace files

When I used the `list.files()` command to list the contents of the `/Users/dan/Rbook/data` directory (in Section 4.4.2), the output referred to a file called `booksales.Rdata`. Let's say I want to load the data from this file into my workspace. The way I do this is with the `load()` function. There are two arguments to this function, but the only one we're interested in is

- `file`. This should be a character string that specifies a path to the file that needs to be loaded. You can use an absolute path or a relative path to do so.

Using the absolute file path, the command would look like this:

```
> load( file = "/Users/dan/Rbook/data/booksales.Rdata" )
```

but this is pretty lengthy. Given that the working directory (remember, we changed the directory at the end of Section 4.4.4) is `/Users/dan/Rbook/data`, I could use a relative file path, like so:

```
> load( file = "../data/booksales.Rdata" )
```

However, my preference is usually to change the working directory first, and *then* load the file. What that would look like is this:

```
> setwd( "../data" )      # move to the data directory
> load( "booksales.Rdata" ) # load the data
```

If I were then to type `who()` I'd see that there are several new variables in my workspace now. Throughout this book, whenever you see me loading a file, I will assume that the file is actually stored in the working directory, or that you've changed the working directory so that R is pointing at the directory that contains the file.

All of the above being said, I feel obliged to point out that Rstudio provides a much easier way to load data files (as long as you're not loading too many files, that is), by using the “file” panel.¹² All you have to do is move to the right directory (see Section 4.4.4), and then click on the name of the file that you want to load. When you do that, Rstudio will attempt to figure out what the right command is to open that kind of file, and then load it. For instance, if I were to click on the `booksales.Rdata` link shown in Figure 4.2, Rstudio would send the following command to the R console for me:

```
> load("~/Rbook/data/booksales.Rdata")
```

Notice that Rstudio specifies the location relative to the home directory `~`.

4.5.2 Importing data from CSV files

One quite commonly used data format is the humble “comma separated value” file, also called a CSV file, and usually bearing the file extension `.csv`. CSV files are just plain old-fashioned text files, and

¹²It also provides an option to “Load Workspace” in the menu system.

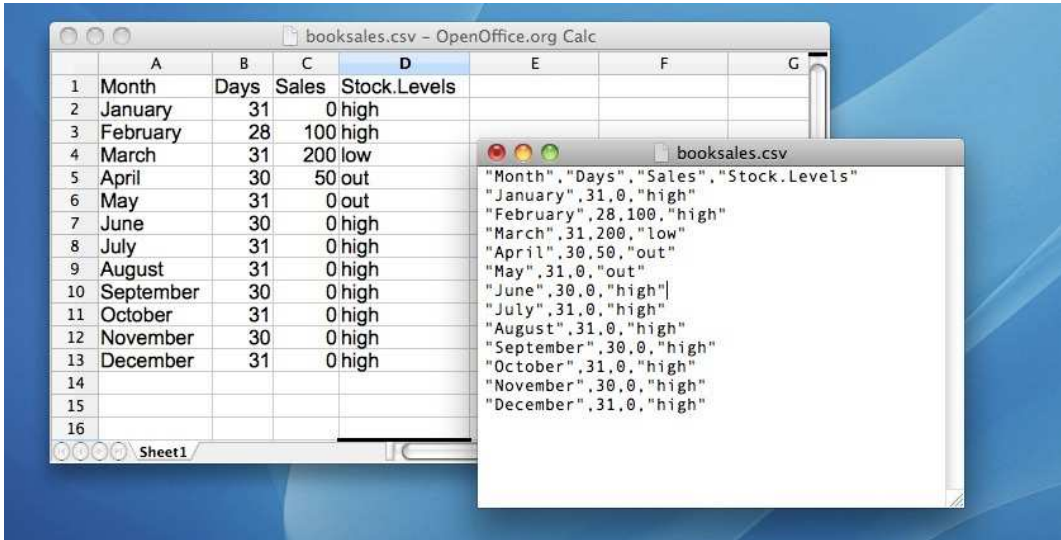


Figure 4.3: The `booksales.csv` data file. On the left, I've opened the file in using a spreadsheet program (OpenOffice), which shows that the file is basically a table. On the right, the same file is open in a standard text editor (the TextEdit program on a Mac), which shows how the file is formatted. The entries in the table are wrapped in quote marks and separated by commas.

.....

what they store is basically just a table of data. This is illustrated in Figure 4.3, which shows a file called `booksales.csv` that I've created. As you can see, each row corresponds to a variable, and each row represents the book sales data for one month. The first row doesn't contain actual data though: it has the names of the variables.

The easiest way to open the file is to use the `read.csv()` function.¹³ This function is pretty flexible, and I'll talk a lot more about it's capabilities in Section 7.9 for more details, but for now there's only two arguments to the function that I'll mention:

- **file**. This should be a character string that specifies a path to the file that needs to be loaded. You can use an absolute path or a relative path to do so.
- **header**. This is a logical value indicating whether or not the first row of the file contains variable names. The default value is `TRUE`.

Therefore, to import the CSV file, the command I need is:

```
> books <- read.csv( file = "booksales.csv" )
```

There are two very important points to notice here. Firstly, notice that I *didn't* try to use the `load()` function, because that function is only meant to be used for `.Rdata` files. If you try to use `load()` on other types of data, you get an error. Secondly, notice that when I imported the CSV file I assigned the

¹³In a lot of books you'll see the `read.table()` function used for this purpose instead of `read.csv()`. They're more or less identical functions, with the same arguments and everything. They differ only in the default values.

result to a variable, which I imaginatively called `books`.¹⁴ Let's have a look at what we've got:

```
> print( books )
   Month Days Sales Stock.Levels
1  January   31     0      high
2 February   28    100      high
3   March   31    200      low
4   April   30     50      out
5    May   31     0      out
6   June   30     0      high
7   July   31     0      high
8  August   31     0      high
9 September   30     0      high
10 October   31     0      high
11 November   30     0      high
12 December   31     0      high
```

Clearly, it's worked, but the format of this output is a bit unfamiliar. We haven't seen anything like this before. What you're looking at is a *data frame*, which is a very important kind of variable in R, and one I'll discuss in Section 4.8. For now, let's just be happy that we imported the data and that it looks about right.

4.5.3 Saving a workspace file

Not surprisingly, saving data is very similar to loading data. To save the entire contents of the workspace to an `.Rdata` file in Rstudio, go to the "Workspace" menu and select the "Save Workspace File..." option. Or equivalently, you can click on the "Save" option in the workspace panel and select the "Save Workspace As..." option. Pretty straightforward, really.¹⁵

Although Rstudio provides a simple way to save files, it's worth understanding the actual commands involved. There are two commands you can use to do this, `save()` and `save.image()`. If you're happy to save *all* of the variables in your workspace into the data file, then you should use `save.image()`. And if you're happy for R to save the file into the current working directory, all you have to do is this:

```
> save.image( file = "myfile.Rdata" )
```

Since `file` is the first argument, you can shorten this to `save.image("myfile.Rdata")`; and if you want to save to a different directory, then (as always) you need to be more explicit about specifying the path to the file, just as we discussed in Section 4.4. Suppose, however, I have several variables in my workspace, and I only want to save some of them. For instance, I might have this as my workspace:

```
> who()
-- Name --  -- Class --  -- Size --
data        data.frame    3 x 2
handy      character      1
junk       numeric        1
```

¹⁴Note that I didn't do this in my earlier example when loading the `.Rdata` file. There's a reason for this. The idea behind an `.Rdata` file is that it stores a whole workspace. So, if you had the ability to look inside the file yourself you'd see that the data file keeps track of all the variables and their names. So when you `load()` the file, R restores all those original names. CSV files are treated differently: as far as R is concerned, the CSV only stores *one* variable, but that variable is big table. So when you import that table into the workspace, R expects *you* to give it a name.

¹⁵A word of warning: what you don't want to do is use the "File" menu. If you look in the "File" menu you will see "Save" and "Save As..." options. Those save the *workspace*, and so they won't produce `.Rdata` files. You use those options to save *scripts*, and so they'll produce `.R` files. We won't get to those until Chapter 8.

I want to save `data` and `handy`, but not `junk`. But I don't want to delete `junk` right now, because I want to use it for something else later on. This is where the `save()` function is useful, since it lets me indicate exactly which variables I want to save. Here is one way I can use the `save` function to solve my problem:

```
> save(data, handy, file = "myfile.Rdata")
```

Importantly, you *must* specify the name of the `file` argument. The reason is that if you don't do so, R will think that "`myfile.Rdata`" is actually a *variable* that you want to save, and you'll get an error message. Finally, I should mention a second way to specify which variables the `save()` function should save, which is to use the `list` argument. You do so like this:

```
> save.me <- c("data", "handy") # the variables to be saved
> save( file = "booksales2.Rdata", list = save.me ) # the command to save them
```

4.5.4 Other things you might want to save

Until now, we've talked mostly about loading and saving *data*.

- *The output.* Sometimes you might also want to keep a copy of all your interactions with R, including everything that you typed in and everything that R did in response. There are some functions that you can use to get R to write its output to a file rather than to print onscreen (e.g., `sink()`), but to be honest, if you do want to save the R output, the easiest thing to do is to use the mouse to select the relevant text in the R console, go to the “Edit” menu in Rstudio and select “Copy”. The output has now been copied to the clipboard. Now open up your favourite text editor or word processing software, and paste it. And you're done. However, this will only save the contents of the console, not the plots you've drawn (assuming you've drawn some). We'll talk about saving images later on.
- *The history.* Another thing that some people like to save is the history. A “history” is just a text file containing the commands that you've typed. You can inspect your history in Rstudio by looking at the history panel, and you can save the contents of the history by clicking on the save button¹⁶. Better yet, you can double click on a command shown in the history panel, and Rstudio will copy it to the R console – which is a handy way of avoiding unnecessary typing.
- *A script.* While it is possible – and sometimes handy – to save the R output as a method for keeping a copy of your statistical analyses, another option that people use a lot (especially when you move beyond simple “toy” analyses) is to write *scripts*. A script is a text file in which you write out all the commands that you want R to run. You can write your script using whatever software you like. In real world data analysis writing scripts is a key skill – and as you become familiar with R you'll probably find that most of what you do involves scripting rather than typing commands at the R prompt. However, you won't need to do much scripting initially, so we'll leave that until Chapter 8.

4.6

Useful things to know about variables

In Chapter 3 I talked a lot about variables, how they're assigned and some of the things you can do with them, but there's a lot of additional complexities. That's not a surprise of course. However, some of

¹⁶That is, the one that looks like an old 3.5 floppy disk – you know, those things that haven't been in use for about 15 years now. I guess it wouldn't be the same to use a picture of a USB stick.

those issues are worth drawing your attention to now. So that's the goal of this section; to cover a few extra topics. As a consequence, this section is basically a bunch of things that I want to briefly mention, but don't really fit in anywhere else. In short, I'll talk about several different issues in this section, which are only loosely connected to one another.

4.6.1 Special values

The first thing I want to mention are some of the “special” values that you might see R produce. Most likely you'll see them in situations where you were expecting a number, but there are quite a few other ways you can encounter them. These values are `Inf`, `NaN`, `NA` and `NULL`. These values can crop up in various different places, and so it's important to understand what they mean.

- *Infinity* (`Inf`). The easiest of the special values to explain is `Inf`, since it corresponds to a value that is infinitely large. You can also have `-Inf`. The easiest way to get `Inf` is to divide a positive number by 0:

```
> 1 / 0  
[1] Inf
```

In most real world data analysis situations, if you're ending up with infinite numbers in your data, then something has gone awry. Hopefully you'll never have to see them.

- *Not a Number* (`NaN`). The special value of `NaN` is short for “not a number”, and it's basically a reserved keyword that means “there isn't a mathematically defined number for this”. If you can remember your high school maths, remember that it is conventional to say that $0/0$ doesn't have a proper answer: mathematicians would say that $0/0$ is *undefined*. R says that it's not a number:

```
> 0 / 0  
[1] NaN
```

Nevertheless, it's still treated as a “numeric” value. To oversimplify, `NaN` corresponds to cases where you asked a proper numerical question that genuinely has *no meaningful answer*.

- *Not available* (`NA`). `NA` indicates that the value that is “supposed” to be stored here is missing. To understand what this means, it helps to recognise that the `NA` value is something that you're most likely to see when analysing data from real world experiments. Sometimes you get equipment failures, or you lose some of the data, or whatever. The point is that some of the information that you were “expecting” to get from your study is just plain missing. Note the difference between `NA` and `NaN`. For `NaN`, we really do know what's supposed to be stored; it's just that it happens to correspond to something like $0/0$ that doesn't make any sense at all. In contrast, `NA` indicates that we actually don't know what was supposed to be there. The information is *missing*.
- *No value* (`NULL`). The `NULL` value takes this “absence” concept even further. It basically asserts that the variable genuinely has no value whatsoever. This is quite different to both `NaN` and `NA`. For `NaN` we actually know what the value is, because it's something insane like $0/0$. For `NA`, we believe that there is supposed to be a value “out there”, but a dog ate our homework and so we don't quite know what it is. But for `NULL` we strongly believe that there is *no value at all*.

4.6.2 Assigning names to vector elements

One thing that is sometimes a little unsatisfying about the way that R prints out a vector is that the elements come out unlabelled. Here's what I mean. Suppose I've got data reporting the quarterly profits for some company. If I just create a no-frills vector, I have to rely on memory to know which element corresponds to which event. That is:

```
> profit <- c( 3.1, 0.1, -1.4, 1.1 )
> profit
[1] 3.1 0.1 -1.4 1.1
```

You can probably guess that the first element corresponds to the first quarter, the second element to the second quarter, and so on, but that's only because I've told you the back story and because this happens to be a very simple example. In general, it can be quite difficult. This is where it can be helpful to assign `names` to each of the elements. Here's how you do it:

```
> names(profit) <- c("Q1", "Q2", "Q3", "Q4")
> profit
  Q1   Q2   Q3   Q4
3.1  0.1 -1.4  1.1
```

This is a slightly odd looking command, admittedly, but it's not too difficult to follow. All we're doing is assigning a vector of labels (character strings) to `names(profit)`. You can always delete the names again by using the command `names(profit) <- NULL`. The important things to notice are that (a) this does make things much easier to read, but (b) the names at the top aren't the "real" data. The *value* of `profit[1]` is still `3.1`; all I've done is added a *name* to `profit[1]` as well. Nevertheless, names aren't purely cosmetic, since R allows you to pull out particular elements of the vector by referring to their names:

```
> profit["Q1"]
  Q1
3.1
```

And if I ever need to pull out the names themselves, then I just type `names(profit)`.

4.6.3 Variable classes

As we've seen, R allows you to store different kinds of data. In particular, the variables we've defined so far have either been character data (text), numeric data, or logical data.¹⁷ It's important that we remember what kind of information each variable stores (and even more important that R remembers) since different kinds of variables allow you to do different things to them. For instance, if your variables have numerical information in them, then it's okay to multiply them together:

```
> x <- 5 # x is numeric
> y <- 4 # y is numeric
> x * y
[1] 20
```

But if they contain character data, multiplication makes no sense whatsoever, and R will complain if you try to do it:

¹⁷Or functions. But let's ignore functions for the moment.

```

> x <- "apples"    # x is character
> y <- "oranges"   # y is character
> x * y
Error in x * y : non-numeric argument to binary operator

```

Even R is smart enough to know you can't multiply "`apples`" by "`oranges`". It knows this because the quote marks are indicators that the variable is supposed to be treated as text, not as a number.

This is quite useful, but notice that it means that R makes a big distinction between `5` and `"5"`. Without quote marks, R treats `5` as the number five, and will allow you to do calculations with it. With the quote marks, R treats `"5"` as the textual character five, and doesn't recognise it as a number any more than it recognises `"p"` or `"five"` as numbers. As a consequence, there's a big difference between typing `x <- 5` and typing `x <- "5"`. In the former, we're storing the number `5`; in the latter, we're storing the character `"5"`. Thus, if we try to do multiplication with the character versions, R gets stroppy:

```

> x <- "5"    # x is character
> y <- "4"    # y is character
> x * y
Error in x * y : non-numeric argument to binary operator

```

Okay, let's suppose that I've forgotten what kind of data I stored in the variable `x` (which happens depressingly often). R provides a function that will let us find out. Or, more precisely, it provides *three* functions: `class()`, `mode()` and `typeof()`. Why the heck does it provide three functions, you might be wondering? Basically, because R actually keeps track of three different kinds of information about a variable:

1. The `class` of a variable is a “high level” classification, and it captures psychologically (or statistically) meaningful distinctions. For instance `"2011-09-12"` and `"my birthday"` are both text strings, but there's an important difference between the two: one of them is a date. So it would be nice if we could get R to recognise that `"2011-09-12"` is a date, and allow us to do things like add or subtract from it. The class of a variable is what R uses to keep track of things like that. Because the class of a variable is critical for determining what R can or can't do with it, the `class()` function is very handy.
2. The `mode` of a variable refers to the format of the information that the variable stores. It tells you whether R has stored text data or numeric data, for instance, which is kind of useful, but it only makes these “simple” distinctions. It can be useful to know about, but it's not the main thing we care about. So I'm not going to use the `mode()` function very much.
3. The `type` of a variable is a very low level classification. We won't use it in this book, but (for those of you that care about these details) this is where you can see the distinction between integer data, double precision numeric, etc. Almost none of you actually will care about this, so I'm not even going to bother demonstrating the `typeof()` function.

For purposes, it's the `class()` of the variable that we care most about. Later on, I'll talk a bit about how you can convince R to “coerce” a variable to change from one class to another (Section 7.10). That's a useful skill for real world data analysis, but it's not something that we need right now. In the meantime, the following examples illustrate the use of the `class()` function:

```

> x <- "hello world"      # x is text
> class(x)
[1] "character"

> x <- TRUE      # x is logical

```

```

> class(x)
[1] "logical"

> x <- 100      # x is a number
> class(x)
[1] "numeric"

```

Exciting, no?

4.7

Factors

Okay, it's time to start introducing some of the data types that are somewhat more specific to statistics. If you remember back to Chapter 2, when we assign numbers to possible outcomes, these numbers can mean quite different things depending on what kind of variable we are attempting to measure. In particular, we commonly make the distinction between *nominal*, *ordinal*, *interval* and *ratio* scale data. How do we capture this distinction in R? Currently, we only seem to have a single numeric data type. That's probably not going to be enough, is it?

A little thought suggests that the numeric variable class in R is perfectly suited for capturing ratio scale data. For instance, if I were to measure response time (RT) for five different events, I could store the data in R like this:

```
> RT <- c(342, 401, 590, 391, 554)
```

where the data here are measured in milliseconds, as is conventional in the psychological literature. It's perfectly sensible to talk about "twice the response time", $2 \times \text{RT}$, or the "response time plus 1 second", $\text{RT} + 1000$, and so both of the following are perfectly reasonable things for R to do:

```

> 2 * RT
[1] 684 802 1180 782 1108
> RT + 1000
[1] 1342 1401 1590 1391 1554

```

And to a lesser extent, the "numeric" class is okay for interval scale data, as long as we remember that multiplication and division aren't terribly interesting for these sorts of variables. That is, if my IQ score is 110 and yours is 120, it's perfectly okay to say that you're 10 IQ points smarter than me¹⁸, but it's not okay to say that I'm only 92% as smart as you are, because intelligence doesn't have a natural zero.¹⁹ We might even be willing to tolerate the use of numeric variables to represent ordinal scale variables, such as those that you typically get when you ask people to rank order items (e.g., like we do in Australian elections), though as we will see R actually has a built in tool for representing ordinal data (see Section 7.11.2) However, when it comes to nominal scale data, it becomes completely unacceptable, because almost all of the "usual" rules for what you're allowed to do with numbers don't apply to nominal scale data. It is for this reason that R has **factors**.

¹⁸Taking all the usual caveats that attach to IQ measurement as a given, of course.

¹⁹Or, more precisely, we don't know how to measure it. Arguably, a rock has zero intelligence. But it doesn't make sense to say that the IQ of a rock is 0 in the same way that we can say that the average human has an IQ of 100. And without knowing what the IQ value is that corresponds to a literal absence of any capacity to think, reason or learn, then we really can't multiply or divide IQ scores and expect a meaningful answer.

4.7.1 Introducing factors

Suppose, I was doing a study in which people could belong to one of three different treatment conditions. Each group of people were asked to complete the same task, but each group received different instructions. Not surprisingly, I might want to have a variable that keeps track of what group people were in. So I could type in something like this

```
> group <- c(1,1,1,2,2,2,3,3,3)
```

so that `group[i]` contains the group membership of the `i`-th person in my study. Clearly, this is numeric data, but equally obviously this is a nominal scale variable. There's no sense in which "group 1" plus "group 2" equals "group 3", but nevertheless if I try to do that, R won't stop me because it doesn't know any better:

```
> group + 2  
[1] 3 3 3 4 4 4 5 5 5
```

Apparently R seems to think that it's allowed to invent "group 4" and "group 5", even though they didn't actually exist. Unfortunately, R is too stupid to know any better: it thinks that `3` is an ordinary number in this context, so it sees no problem in calculating `3 + 2`. But since we're not that stupid, we'd like to stop R from doing this. We can do so by instructing R to treat `group` as a factor. This is easy to do using the `as.factor()` function.²⁰

```
> group <- as.factor(group)  
> group  
[1] 1 1 1 2 2 2 3 3 3  
Levels: 1 2 3
```

It looks more or less the same as before (though it's not immediately obvious what all that `Levels` rubbish is about), but if we ask R to tell us what the class of the `group` variable is now, it's clear that it has done what we asked:

```
> class(group)  
[1] "factor"
```

Neat. Better yet, now that I've converted `group` to a factor, look what happens when I try to add 2 to it:

```
> group + 2  
[1] NA NA NA NA NA NA NA NA NA  
Warning message:  
In Ops.factor(group, 2) : + not meaningful for factors
```

This time even R is smart enough to know that I'm being an idiot, so it tells me off and then produces a vector of missing values.

4.7.2 Labelling the factor levels

I have a confession to make. My memory is not infinite in capacity; and it seems to be getting worse as I get older. So it kind of annoys me when I get data sets where there's a nominal scale variable called

²⁰Once again, this is an example of *coercing* a variable from one class to another. I'll talk about coercion in more detail in Section 7.10.

`gender`, with two levels corresponding to males and females. But when I go to print out the variable I get something like this:

```
> gender  
[1] 1 1 1 1 2 2 2  
Levels: 1 2
```

Okaaaay. That's not helpful at all, and it makes me very sad. Which number corresponds to the males and which one corresponds to the females? Wouldn't it be nice if R could actually keep track of this? It's way too hard to remember which number corresponds to which gender. And besides, the problem that this causes is much more serious than a single sad nerd... because R has no way of knowing that the `1`s in the `group` variable are a very different kind of thing to the `1`s in the `gender` variable. So if I try to ask which elements of the `group` variable are equal to the corresponding elements in `gender`, R thinks this is totally kosher, and gives me this:

```
> group == gender  
[1] TRUE TRUE TRUE FALSE FALSE TRUE FALSE FALSE FALSE
```

Well, that's ... especially stupid.²¹ The problem here is that R is very literal minded. Even though you've declared both `group` and `gender` to be factors, it still assumes that a `1` is a `1` no matter which variable it appears in.

To fix both of these problems (my memory problem, and R's infuriating literal interpretations), what we need to do is assign meaningful labels to the different *levels* of each factor. We can do that like this:

```
> levels(group) <- c("group 1", "group 2", "group 3")  
> print(group)  
[1] group 1 group 1 group 1 group 2 group 2 group 2 group 3 group 3 group 3  
Levels: group 1 group 2 group 3  
>  
> levels(gender) <- c("male", "female")  
> print(gender)  
[1] male male male male female female female female  
Levels: male female
```

That's much easier on the eye, and better yet, R is smart enough to know that "`female`" is not equal to "`group 2`", so now when I try to ask which group memberships are "equal to" the gender of the corresponding person,

```
> group == gender  
Error in Ops.factor(group, gender) : level sets of factors are different
```

R correctly tells me that I'm an idiot.

4.7.3 Moving on...

Factors are very useful things, and we'll use them a lot in this book: they're *the* main way to represent a nominal scale variable. And there are lots of nominal scale variables out there. I'll talk more about

²¹Some users might wonder why R even allows the `==` operator for factors. The reason is that sometimes you really do have different factors that have the same levels. For instance, if I was analysing data associated with football games, I might have a factor called `home.team`, and another factor called `winning.team`. In that situation I really should be able to ask if `home.team == winning.team`.

factors in 7.11.2, but for now you know enough to be able to get started.

4.8

Data frames

It's now time to go back and deal with the somewhat confusing thing that happened in Section 4.5.2 when we tried to open up a CSV file. Apparently we succeeded in loading the data, but it came to us in a very odd looking format. At the time, I told you that this was a **data frame**. Now I'd better explain what that means.

4.8.1 Introducing data frames

In order to understand why R has created this funny thing called a data frame, it helps to try to see what problem it solves. So let's go back to the little scenario that I used when introducing factors in Section 4.7. In that section I recorded the **group** and **gender** for all 9 participants in my study. Let's also suppose I recorded their ages and their **score** on "Dan's Terribly Exciting Psychological Test":

```
> age <- c(17, 19, 21, 37, 18, 19, 47, 18, 19)
> score <- c(12, 10, 11, 15, 16, 14, 25, 21, 29)
```

Assuming no other variables are in the workspace, if I type **who()** I get this:

```
> who()
-- Name --  -- Class --  -- Size --
age        numeric      9
gender     factor       9
group      factor       9
score      numeric      9
```

So there are four variables in the workspace, **age**, **gender**, **group** and **score**. And it just so happens that all four of them are the same size (i.e., they're all vectors with 9 elements). Aaaand it just so happens that **age[1]** corresponds to the age of the first person, and **gender[1]** is the gender of that very same person, etc. In other words, you and I both know that all four of these variables correspond to the *same* data set, and all four of them are organised in exactly the same way.

However, R *doesn't* know this! As far as it's concerned, there's no reason why the **age** variable has to be the same length as the **gender** variable; and there's no particular reason to think that **age[1]** has any special relationship to **gender[1]** any more than it has a special relationship to **gender[4]**. In other words, when we store everything in separate variables like this, R doesn't know anything about the relationships between things. It doesn't even really know that these variables actually refer to a proper data set. The data frame fixes this: if we store our variables inside a data frame, we're telling R to treat these variables as a single, fairly coherent data set.

To see how they do this, let's create one. So how do we create a data frame? One way we've already seen: if we import our data from a CSV file, R will store it as a data frame. A second way is to create it directly from some existing variables using the **data.frame()** function. All you have to do is type a list of variables that you want to include in the data frame. The output of a **data.frame()** command is, well, a data frame. So, if I want to store all four variables from my experiment in a data frame called **expt** I can do so like this:

```

> expt <- data.frame ( age, gender, group, score )
> expt
  age gender group score
1 17 male group 1 12
2 19 male group 1 10
3 21 male group 1 11
4 37 male group 2 15
5 18 male group 2 16
6 19 female group 2 14
7 47 female group 3 25
8 18 female group 3 21
9 19 female group 3 29

```

Note that `expt` is a completely self-contained variable. Once you've created it, it no longer depends on the original variables from which it was constructed. That is, if we make changes to the original `age` variable, it will *not* lead to any changes to the age data stored in `expt`.

4.8.2 Pulling out the contents of the data frame using `$`

At this point, our workspace contains only the one variable, a data frame called `expt`. But as we can see when we told R to print the variable out, this data frame contains 4 variables, each of which has 9 observations. So how do we get this information out again? After all, there's no point in storing information if you don't use it, and there's no way to use information if you can't access it. So let's talk a bit about how to pull information out of a data frame.

The first thing we might want to do is pull out one of our stored variables, let's say `score`. One thing you might try to do is ignore the fact that `score` is locked up inside the `expt` data frame. For instance, you might try to print it out like this:

```

> score
Error: object 'score' not found

```

This doesn't work, because R doesn't go "peeking" inside the data frame unless you explicitly tell it to do so. There's actually a very good reason for this, which I'll explain in a moment, but for now let's just assume R knows what it's doing. How do we tell R to look inside the data frame? As is always the case with R there are several ways. The simplest way is to use the `$` operator to extract the variable you're interested in, like this:

```

> expt$score
[1] 12 10 11 15 16 14 25 21 29

```

4.8.3 Pulling out the contents of the data frame using `with()`

If all you want to do is extract one of the stored variables, you'll probably find that the `$` operator is the easiest thing to do. But there are other possibilities too. One alternative is to use the `with()` function. The `with()` function has two arguments:

- `data`. This is the name of the data frame.
- `expr`. This can be any expression (i.e., command) that you want evaluated. It will be evaluated using the variables stored in the data frame.

So if you wanted to print out the `score` variable, you could use the `with()` function to do so. First, here's how it looks if I specify both arguments by name:

```
> with( data = expt,           # the data frame is 'expt'  
+       expr = print(score)  # what I want is to print out 'score'  
+ )  
[1] 12 10 11 15 16 14 25 21 29
```

Clearly, this is just an extremely long and tedious way of doing the exact same thing that `expt$score` does, so there's really no value to doing it this way. Of course, you can shorten it to this

```
> with( expt, score )  
[1] 12 10 11 15 16 14 25 21 29
```

but that's not a huge improvement. It's still awfully long.

Where the `with()` function starts to become useful is when you need to do something more complicated using the variables in the data frame. To take a simple example, suppose you want to R to print out the scores of the teenaged males. If we use `$` to do the work, here's what the command looks like:

```
> expt$score[ expt$age < 20 &  
+             expt$gender == "male"  
+ ]  
[1] 12 10 16
```

where I've split the command over multiple lines to try to make it look as clean as possible. Alternatively, I could use the `with()` function. Again, I split the command over multiple lines to make it look nice, which looks like this:

```
> with( expt,  
+       score[ age < 20 &  
+                 gender == "male"  
+               ]  
+ )  
[1] 12 10 16
```

The two versions involve a similar number of characters, but (at least to my eye) the version that uses `with()` is slightly easier to read.

4.8.4 Getting information about a data frame

One problem that sometimes comes up in practice is that you forget what you called all your variables. Normally you might try to type `objects()` or `who()`, but neither of those commands will tell you what the names are for those variables inside a data frame! One way is to ask R to tell you what the *names* of all the variables stored in the data frame are, which you can do using the `names()` function:

```
> names(expt)  
[1] "age"    "gender"  "group"   "score"
```

An alternative method is to use the `who()` function, as long as you tell it to look at the variables inside data frames. If you set `expand = TRUE` then it will not only list the variables in the workspace, but it will “expand” any data frames that you've got in the workspace, so that you can see what they look like. That is:

```
> who(expand = TRUE)
-- Name --  -- Class --  -- Size --
expt      data.frame   9 x 4
$age      numeric      9
$gender   factor       9
$group    factor       9
$score    numeric      9
```

or, since `expand` is the first argument in the `who()` function you can just type `who(TRUE)` or even `who(T)` instead. I'll do that a lot in this book.

4.8.5 Looking for more on data frames?

There's a lot more that can be said about data frames: they're fairly complicated beasts, and the longer you use R the more important it is to make sure you really understand them. We'll talk a lot more about them in Chapter 7.

4.9 _____

Lists

The next kind of data I want to mention are **lists**. Lists are an extremely fundamental data structure in R, and as you start making the transition from a novice to a savvy R user you will use lists all the time. I don't use lists very often in this book – not directly – but most of the advanced data structures in R are built from lists (e.g., data frames are actually a specific type of list). Because lists are so important to how R stores things, it's useful to have a basic understanding of them. Okay, so what is a list, exactly? Like data frames, lists are just “collections of variables.” However, unlike data frames – which are basically supposed to look like a nice “rectangular” table of data – there are no constraints on what kinds of variables we include, and no requirement that the variables have any particular relationship to one another. In order to understand what this actually *means*, the best thing to do is create a list, which we can do using the `list()` function. If I type this as my command:

```
> Dan <- list( age = 34,
+               nerd = TRUE,
+               parents = c("Joe", "Liz")
+ )
```

R creates a new list variable called `Dan`, which is a bundle of three different variables: `age`, `nerd` and `parents`. Notice, that the `parents` variable is longer than the others. This is perfectly acceptable for a list, but it wouldn't be for a data frame. If we now print out the variable, you can see the way that R stores the list:

```
> print( Dan )
$age
[1] 34

$nerd
[1] TRUE

$parents
[1] "Joe" "Liz"
```

As you might have guessed from those `$` symbols everywhere, the variables are stored in exactly the same way that they are for a data frame (again, this is not surprising: data frames *are* a type of list). So you will (I hope) be entirely unsurprised and probably quite bored when I tell you that you can extract the variables from the list using the `$` operator, like so:

```
> Dan$nerd  
[1] TRUE
```

If you need to add new entries to the list, the easiest way to do so is to again use `$`, as the following example illustrates. If I type a command like this

```
> Dan$children <- "Alex"
```

then R creates a new entry to the end of the list called `children`, and assigns it a value of `"Alex"`. If I were now to `print()` this list out, you'd see a new entry at the bottom of the printout. Finally, it's actually possible for lists to contain other lists, so it's quite possible that I would end up using a command like `Dan$children$age` to find out how old my son is. Or I could try to remember it myself I suppose.

4.10 _____

Formulas

The last kind of variable that I want to introduce before finally being able to start talking about statistics is the **formula**. Formulas were originally introduced into R as a convenient way to specify a particular type of statistical model (see Chapter 15) but they're such handy things that they've spread. Formulas are now used in a lot of different contexts, so it makes sense to introduce them early.

Stated simply, a formula object is a variable, but it's a special type of variable that specifies a relationship between other variables. A formula is specified using the “tilde operator” `~`. A very simple example of a formula is shown below:²²

```
> formula1 <- out ~ pred  
> formula1  
out ~ pred
```

The *precise* meaning of this formula depends on exactly what you want to do with it, but in broad terms it means “the `out` (outcome) variable, analysed in terms of the `pred` (predictor) variable”. That said, although the simplest and most common form of a formula uses the “one variable on the left, one variable on the right” format, there are others. For instance, the following examples are all reasonably common

```
> formula2 <- out ~ pred1 + pred2    # more than one variable on the right  
> formula3 <- out ~ pred1 * pred2    # different relationship between predictors  
> formula4 <- ~ var1 + var2        # a 'one-sided' formula
```

and there are many more variants besides. Formulas are pretty flexible things, and so different functions will make use of different formats, depending on what the function is intended to do.

²²Note that, when I write out the formula, R doesn't check to see if the `out` and `pred` variables actually exist: it's only later on when you try to use the formula for something that this happens.

4.11

Generic functions

There's one really important thing that I omitted when I discussed functions earlier on in Section 3.5, and that's the concept of a **generic function**. The two most notable examples that you'll see in the next few chapters are `summary()` and `plot()`, although you've already seen an example of one working behind the scenes, and that's the `print()` function. The thing that makes generics different from the other functions is that their behaviour changes, often quite dramatically, depending on the `class()` of the input you give it. The easiest way to explain the concept is with an example. With that in mind, let's take a closer look at what the `print()` function actually does. I'll do this by creating a formula, and printing it out in a few different ways. First, let's stick with what we know:

```
> my.formula <- blah ~ blah.blah      # create a variable of class "formula"
> print( my.formula )                # print it out using the generic print() function
blah ~ blah.blah
```

So far, there's nothing very surprising here. But there's actually a lot going on behind the scenes here. When I type `print(my.formula)`, what actually happens is the `print()` function checks the class of the `my.formula` variable. When the function discovers that the variable it's been given is a formula, it goes looking for a function called `print.formula()`, and then delegates the whole business of printing out the variable to the `print.formula()` function.²³ For what it's worth, the name for a “dedicated” function like `print.formula()` that exists only to be a special case of a generic function like `print()` is a **method**, and the name for the process in which the generic function passes off all the hard work onto a method is called **method dispatch**. You won't need to understand the details at all for this book, but you do need to know the gist of it; if only because a lot of the functions we'll use are actually generics. Anyway, to help expose a little more of the workings to you, let's bypass the `print()` function entirely and call the formula method directly:

```
> print.formula( my.formula )        # print it out using the print.formula() method
blah ~ blah.blah
```

There's no difference in the output at all. But this shouldn't surprise you because it was actually the `print.formula()` method that was doing all the hard work in the first place. The `print()` function itself is a lazy bastard that doesn't do anything other than select which of the methods is going to do the actual printing.

Okay, fair enough, but you might be wondering what would have happened if `print.formula()` didn't exist? That is, what happens if there isn't a specific method defined for the class of variable that you're using? In that case, the generic function passes off the hard work to a “default” method, whose name in this case would be `print.default()`. Let's see what happens if we bypass the `print()` formula, and try to print out `my.formula` using the `print.default()` function:

```
> print.default( my.formula )       # print it out using the print.default() method
blah ~ blah.blah
attr(,"class")
[1] "formula"
attr(,".Environment")
```

²³For readers with a programming background: what I'm describing is the very basics of how S3 methods work. However, you should be aware that R has two entirely distinct systems for doing object oriented programming, known as S3 and S4. Of the two, S3 is simpler and more informal, whereas S4 supports all the stuff that you might expect of a fully object oriented language. Most of the generics we'll run into in this book use the S3 system, which is convenient for me because I'm still trying to figure out S4.

```
<environment: R_GlobalEnv>
<environment: R_GlobalEnv>
```

Hm. You can kind of see that it is trying to print out the same formula, but there's a bunch of ugly low-level details that have also turned up on screen. This is because the `print.default()` method doesn't know anything about formulas, and doesn't know that it's supposed to be hiding the obnoxious internal gibberish that R produces sometimes.

At this stage, this is about as much as we need to know about generic functions and their methods. In fact, you can get through the entire book without learning any more about them than this, so it's probably a good idea to end this discussion here.

4.12

Getting help

The very last topic I want to mention in this chapter is where to go to find help. Obviously, I've tried to make this book as helpful as possible, but it's not even close to being a comprehensive guide, and there's thousands of things it doesn't cover. So where should you go for help?

4.12.1 How to read the help documentation

I have somewhat mixed feelings about the help documentation in R. On the plus side, there's a lot of it, and it's very thorough. On the minus side, there's a lot of it, and it's very thorough. There's so much help documentation that it sometimes doesn't help, and most of it is written with an advanced user in mind. Often it feels like most of the help files work on the assumption that the reader already understands everything about R except for the specific topic that it's providing help for. What that means is that, once you've been using R for a long time and are beginning to get a feel for how to use it, the help documentation is *awesome*. These days, I find myself really liking the help files (most of them anyway). But when I first started using R I found it very dense.

To some extent, there's not much I can do to help you with this. You just have to work at it yourself; once you're moving away from being a pure beginner and are becoming a skilled user, you'll start finding the help documentation more and more helpful. In the meantime, I'll help as much as I can by trying to explain to you what you're looking at when you open a help file. To that end, let's look at the help documentation for the `load()` function. To do so, I type either of the following:

```
> ?load
> help("load")
```

When I do that, R goes looking for the help file for the "`load`" topic. If it finds one, Rstudio takes it and displays it in the help panel. Alternatively, you can try a fuzzy search for a help topic:

```
> ??load
> help.search("load")
```

This will bring up a list of possible topics that you might want to follow up in.

Regardless, at some point you'll find yourself looking at an actual help file. And when you do, you'll see there's a quite a lot of stuff written down there, and it comes in a pretty standardised format. So let's go through it slowly, using the "`load`" topic as our example. Firstly, at the very top we see this:

The R Documentation part isn't terribly interesting, but the `load { base }` part is telling you that this is in reference to the `load()` function (obviously) and that this function is in the `base` package. Next, we get the “title” information, and a short “description” of what the function does:

Reload Saved Datasets

Description

Reload datasets written with the function `save`.

Fairly straightforward. The next section describes how the function is used:

Usage

```
load(file, envir = parent.frame())
```

In this instance, the usage section is actually pretty readable. It's telling you that there are two arguments to the `load()` function: the first one is called `file`, and the second one is called `envir`. It's also telling you that there is a default value for the `envir` argument; so if the user doesn't specify what the value of `envir` should be, then R will assume that `envir = parent.frame()`. In contrast, the `file` argument has no default value at all, so the user *must* specify a value for it. So in one sense, this section is very straightforward.

The problem, of course, is that you don't know what the `parent.frame()` function actually does, so it's hard for you to know what the `envir = parent.frame()` bit is all about. What you *could* do is then go look up the help documents for the `parent.frame()` function (and sometimes that's actually a good idea), but often you'll find that the help documents for those functions are just as dense (if not more dense) than the help file that you're currently reading. As an alternative, my general approach when faced with something like this is to skim over it, see if I can make any sense of it. If so, great. If not, I find that the best thing to do is ignore it. In fact, the first time I read the help file for the `load()` function, I had no idea what any of the `envir` related stuff was about. But fortunately I didn't have to: the default setting here (i.e., `envir = parent.frame()`) is actually the thing you want in about 99% of cases, so it's safe to ignore it.

Basically, what I'm trying to say is: don't let the scary, incomprehensible parts of the help file intimidate you. Especially because there's often some parts of the help file that will make sense. Of course, I guarantee you that sometimes this strategy will lead you to make mistakes... often embarrassing mistakes. But it's still better than getting paralysed with fear.

So, let's continue on. The next part of the help documentation discusses each of the arguments, and what they're supposed to do:

Arguments

- | | |
|--------------------|---|
| <code>file</code> | a (readable binary-mode) connection or a character string giving the name of the file to load (when tilde expansion is done). |
| <code>envir</code> | the environment where the data should be loaded. |

Okay, so what this is telling us is that the `file` argument needs to be a string (i.e., text data) which tells R the name of the file to load. It also seems to be hinting that there's other possibilities too (e.g., a "binary mode connection"), and you probably aren't quite sure what "tilde expansion" means²⁴. But overall, the meaning is pretty clear.

Turning to the `envir` argument, it's now a little clearer what the Usage section was babbling about. The `envir` argument specifies the name of an environment (see Section 4.3 if you've forgotten what environments are) into which R should place the variables when it loads the file. Almost always, this is a no-brainer: you want R to load the data into the same damn environment in which you're invoking the `load()` command. That is, if you're typing `load()` at the R prompt, then you want the data to be loaded into your workspace (i.e., the global environment). But if you're writing your own function that needs to load some data, you want the data to be loaded inside that function's private workspace. And in fact, that's exactly what the `parent.frame()` thing is all about. It's telling the `load()` function to send the data to the same place that the `load()` command itself was coming from. As it turns out, if we'd just ignored the `envir` bit we would have been totally safe. Which is nice to know.

Moving on, next up we get a detailed description of what the function actually does:

Details

`load` can load R objects saved in the current or any earlier format. It can read a compressed file (see `save`) directly from a file or from a suitable connection (including a call to `url`).

A not-open connection will be opened in mode "`rb`" and closed after use. Any connection other than a `gzfile` or `gzcon` connection will be wrapped in `gzcon` to allow compressed saves to be handled: note that this leaves the connection in an altered state (in particular, binary-only).

Only R objects saved in the current format (used since R 1.4.0) can be read from a connection. If no input is available on a connection a warning will be given, but any input not in the current format will result in a error.

Loading from an earlier version will give a warning about the "magic number": magic numbers `1971:1977` are from R < 0.99.0, and `RD[ABX]1` from R 0.99.0 to R 1.3.1. These are all obsolete, and you are strongly recommended to re-save such files in a current format.

Then it tells you what the output value of the function is:

Value

A character vector of the names of objects created, invisibly.

This is usually a bit more interesting, but since the `load()` function is mainly used to load variables into the workspace rather than to return a value, it's no surprise that this doesn't do much or say much. Moving on, we sometimes see a few additional sections in the help file, which can be different depending on what the function is:

²⁴It's extremely simple, by the way. We discussed it in Section 4.4, though I didn't call it by that name. Tilde expansion is the thing where R recognises that, in the context of specifying a file location, the tilde symbol `~` corresponds to the user home directory (e.g., `/Users/dan/`).

Warning

Saved R objects are binary files, even those saved with `ascii = TRUE`, so ensure that they are transferred without conversion of end of line markers. `load` tries to detect such a conversion and gives an informative error message.

Yeah, yeah. Warning, warning, blah blah blah. Towards the bottom of the help file, we see something like this, which suggests a bunch of related topics that you might want to look at. These can be quite helpful:

See Also

`save`, `download.file`.

For other interfaces to the underlying serialization format, see `unserialize` and `readRDS`.

Finally, it gives you some examples of how to use the function(s) that the help file describes. These are supposed to be proper R commands, meaning that you should be able to type them into the console yourself and they'll actually work. Sometimes it can be quite helpful to try the examples yourself. Anyway, here they are for the "`load`" help file:

Examples

```
## save all data
xx <- pi # to ensure there is some data
save(list = ls(all=TRUE), file= "all.RData")
rm(xx)

## restore the saved values to the current environment
local({
  load("all.RData")
  ls()
})
## restore the saved values to the user's workspace
load("all.RData", .GlobalEnv)

unlink("all.RData")

## Not run:
con <- url("http://some.where.net/R/data/example.rda")
## print the value to see what objects were created.
print(load(con))
close(con) # url() always opens the connection

## End(Not run)
```

As you can see, they're pretty dense, and not at all obvious to the novice user. However, they do provide good examples of the various different things that you can do with the `load()` function, so it's not a bad idea to have a look at them, and to try not to find them too intimidating.

4.12.2 Other resources

- The Rseek website (www.rseek.org). One thing that I really find annoying about the R help documentation is that it's hard to search properly. When coupled with the fact that the documentation

is dense and highly technical, it's often a better idea to search or ask online for answers to your questions. With that in mind, the Rseek website is great: it's an R specific search engine. I find it really useful, and it's almost always my first port of call when I'm looking around.

- The R-help mailing list (see <http://www.r-project.org/mail.html> for details). This is the official R help mailing list. It can be very helpful, but it's *very* important that you do your homework before posting a question. The list gets a lot of traffic. While the people on the list try as hard as they can to answer questions, they do so for free, and you *really* don't want to know how much money they could charge on an hourly rate if they wanted to apply market rates. In short, they are doing you a favour, so be polite. Don't waste their time asking questions that can be easily answered by a quick search on Rseek (it's rude), make sure your question is clear, and all of the relevant information is included. In short, read the posting guidelines carefully (<http://www.r-project.org/posting-guide.html>), and make use of the `help.request()` function that R provides to check that you're actually doing what you're expected.

4.13

Summary

- Installing, loading and updating packages (Section 4.2)
- Viewing the workspace and removing variables (Section 4.3)
- Getting around your computer (Section 4.4)
- Loading and saving data (Section 4.5)
- Useful things to know about variables: special values, element names and classes (Section 4.6)
- More complex types of variables: factors (Section 4.7), data frames (Section 4.8), lists (Section 4.9) and formulas (Section 4.10).
- Generic functions (Section 4.11)
- Getting help (Section 4.12)

Part III.

Working with data

5. Descriptive statistics

Any time that you get a new data set to look at, one of the first tasks that you have to do is find ways of summarising the data in a compact, easily understood fashion. This is what **descriptive statistics** (as opposed to inferential statistics) is all about. In fact, to many people the term “statistics” is synonymous with descriptive statistics. It is this topic that we’ll consider in this chapter, but before going into any details, let’s take a moment to get a sense of why we need descriptive statistics. To do this, let’s load the `aflsmall.Rdata` file, and use the `who()` function in the `lsr` package to see what variables are stored in the file:

```
> load("aflsmall.Rdata")
> library(lsr)
> who()
-- Name --      -- Class --      -- Size --
afl.finalists   factor          400
afl.margins     numeric         176
```

There are two variables here, `afl.finalists` and `afl.margins`. We’ll focus a bit on these two variables in this chapter, so I’d better tell you what they are. Unlike most of data sets in this book, these are actually real data, relating to the Australian Football League (AFL)¹. The `afl.margins` variable contains the winning margin (number of points) for all 176 home and away games played during the 2010 season. The `afl.finalists` variable contains the names of all 400 teams that played in all 200 finals matches played during the period 1987 to 2010. Let’s have a look at the `afl.margins` variable:

```
> print(afl.margins)
 [1]  56  31  56   8  32  14  36  56  19   1   3 104  43  44  72   9  28
[18]  25  27  55  20  16  16   7  23  40  48  64  22  55  95  15  49  52
[35]  50  10  65  12  39  36   3  26  23  20  43 108  53  38   4   8   3
[52]  13  66  67  50  61  36  38  29   9  81   3  26  12  36  37  70   1
[69]  35  12  50  35   9  54  47   8  47   2  29  61  38  41  23  24   1
[86]   9  11  10  29  47  71  38  49  65  18   0  16   9  19  36  60  24
[103]  25  44  55   3  57  83  84  35   4  35  26  22   2  14  19  30  19
[120]  68  11  75  48  32  36  39  50  11   0  63  82  26   3  82  73  19
[137]  33  48   8  10  53  20  71  75  76  54  44   5  22  94  29   8  98
[154]   9  89   1 101   7  21  52  42  21 116   3  44  29  27  16   6  44
[171]   3  28  38  29  10   10
```

This output doesn’t make it easy to get a sense of what the data are actually saying. Just “looking at the data” isn’t a terribly effective way of understanding data. In order to get some idea about what’s going on, we need to calculate some descriptive statistics (this chapter) and draw some nice pictures

¹Note for non-Australians: the AFL is an Australian rules football competition. You don’t need to know anything about Australian rules in order to follow this section.

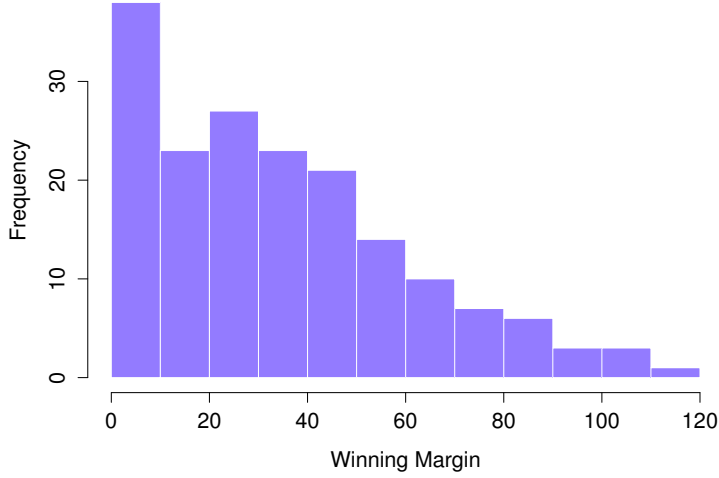


Figure 5.1: A histogram of the AFL 2010 winning margin data (the `afl.margins` variable). As you might expect, the larger the margin the less frequently you tend to see it.

.....

(Chapter 6). Since the descriptive statistics are the easier of the two topics, I'll start with those, but nevertheless I'll show you a histogram of the `afl.margins` data, since it should help you get a sense of what the data we're trying to describe actually look like. But for what it's worth, this histogram – which is shown in Figure 5.1 – was generated using the `hist()` function. We'll talk a lot more about how to draw histograms in Section 6.3. For now, it's enough to look at the histogram and note that it provides a fairly interpretable representation of the `afl.margins` data.

5.1 _____

Measures of central tendency

Drawing pictures of the data, as I did in Figure 5.1 is an excellent way to convey the “gist” of what the data is trying to tell you, it’s often extremely useful to try to condense the data into a few simple “summary” statistics. In most situations, the first thing that you’ll want to calculate is a measure of **central tendency**. That is, you’d like to know something about the “average” or “middle” of your data lies. The two most commonly used measures are the mean, median and mode; occasionally people will also report a trimmed mean. I’ll explain each of these in turn, and then discuss when each of them is useful.

5.1.1 The mean

The **mean** of a set of observations is just a normal, old-fashioned average: add all of the values up, and then divide by the total number of values. The first five AFL margins were 56, 31, 56, 8 and 32, so the

mean of these observations is just:

$$\frac{56 + 31 + 56 + 8 + 32}{5} = \frac{183}{5} = 36.60$$

Of course, this definition of the mean isn't news to anyone: averages (i.e., means) are used so often in everyday life that this is pretty familiar stuff. However, since the concept of a mean is something that everyone already understands, I'll use this as an excuse to start introducing some of the mathematical notation that statisticians use to describe this calculation, and talk about how the calculations would be done in R.

The first piece of notation to introduce is N , which we'll use to refer to the number of observations that we're averaging (in this case $N = 5$). Next, we need to attach a label to the observations themselves. It's traditional to use X for this, and to use subscripts to indicate which observation we're actually talking about. That is, we'll use X_1 to refer to the first observation, X_2 to refer to the second observation, and so on, all the way up to X_N for the last one. Or, to say the same thing in a slightly more abstract way, we use X_i to refer to the i -th observation. Just to make sure we're clear on the notation, the following table lists the 5 observations in the `afl.margins` variable, along with the mathematical symbol used to refer to it, and the actual value that the observation corresponds to:

the observation	its symbol	the observed value
winning margin, game 1	X_1	56 points
winning margin, game 2	X_2	31 points
winning margin, game 3	X_3	56 points
winning margin, game 4	X_4	8 points
winning margin, game 5	X_5	32 points

Okay, now let's try to write a formula for the mean. By tradition, we use \bar{X} as the notation for the mean. So the calculation for the mean could be expressed using the following formula:

$$\bar{X} = \frac{X_1 + X_2 + \dots + X_{N-1} + X_N}{N}$$

This formula is entirely correct, but it's terribly long, so we make use of the **summation symbol** Σ to shorten it.² If I want to add up the first five observations, I could write out the sum the long way, $X_1 + X_2 + X_3 + X_4 + X_5$ or I could use the summation symbol to shorten it to this:

$$\sum_{i=1}^5 X_i$$

Taken literally, this could be read as "the sum, taken over all i values from 1 to 5, of the value X_i ". But basically, what it means is "add up the first five observations". In any case, we can use this notation to write out the formula for the mean, which looks like this:

$$\bar{X} = \frac{1}{N} \sum_{i=1}^N X_i$$

In all honesty, I can't imagine that all this mathematical notation helps clarify the concept of the mean at all. In fact, it's really just a fancy way of writing out the same thing I said in words: add all the

²The choice to use Σ to denote summation isn't arbitrary: it's the Greek upper case letter sigma, which is the analogue of the letter S in that alphabet. Similarly, there's an equivalent symbol used to denote the multiplication of lots of numbers: because multiplications are also called "products", we use the Π symbol for this; the Greek upper case pi, which is the analogue of the letter P.

values up, and then divide by the total number of items. However, that's not really the reason I went into all that detail. My goal was to try to make sure that everyone reading this book is clear on the notation that we'll be using throughout the book: \bar{X} for the mean, Σ for the idea of summation, X_i for the i th observation, and N for the total number of observations. We're going to be re-using these symbols a fair bit, so it's important that you understand them well enough to be able to "read" the equations, and to be able to see that it's just saying "add up lots of things and then divide by another thing".

5.1.2 Calculating the mean in R

Okay that's the maths, how do we get the magic computing box to do the work for us? If you really wanted to, you could do this calculation directly in R. For the first 5 AFL scores, do this just by typing it in as if R were a calculator...

```
> (56 + 31 + 56 + 8 + 32) / 5
[1] 36.6
```

...in which case R outputs the answer 36.6, just as if it were a calculator. However, that's not the only way to do the calculations, and when the number of observations starts to become large, it's easily the most tedious. Besides, in almost every real world scenario, you've already got the actual numbers stored in a variable of some kind, just like we have with the `afl.margins` variable. Under those circumstances, what you want is a function that will just add up all the values stored in a numeric vector. That's what the `sum()` function does. If we want to add up all 176 winning margins in the data set, we can do so using the following command:³

```
> sum( afl.margins )
[1] 6213
```

If we only want the sum of the first five observations, then we can use square brackets to pull out only the first five elements of the vector. So the command would now be:

```
> sum( afl.margins[1:5] )
[1] 183
```

To calculate the mean, we now tell R to divide the output of this summation by five, so the command that we need to type now becomes the following:

```
> sum( afl.margins[1:5] ) / 5
[1] 36.6
```

Although it's pretty easy to calculate the mean using the `sum()` function, we can do it in an even easier way, since R also provides us with the `mean()` function. To calculate the mean for all 176 games, we would use the following command:

```
> mean( x = afl.margins )
[1] 35.30114
```

However, since `x` is the first argument to the function, I could have omitted the argument name. In any case, just to show you that there's nothing funny going on, here's what we would do to calculate the mean for the first five observations:

³Note that, just as we saw with the combine function `c()` and the remove function `rm()`, the `sum()` function has unnamed arguments. I'll talk about unnamed arguments later in Section 8.4.1, but for now let's just ignore this detail.

```
> mean( afl.margins[1:5] )
[1] 36.6
```

As you can see, this gives exactly the same answers as the previous calculations.

5.1.3 The median

The second measure of central tendency that people use a lot is the **median**, and it's even easier to describe than the mean. The median of a set of observations is just the middle value. As before let's imagine we were interested only in the first 5 AFL winning margins: 56, 31, 56, 8 and 32. To figure out the median, we sort these numbers into ascending order:

8, 31, **32**, 56, 56

From inspection, it's obvious that the median value of these 5 observations is 32, since that's the middle one in the sorted list (I've put it in bold to make it even more obvious). Easy stuff. But what should we do if we were interested in the first 6 games rather than the first 5? Since the sixth game in the season had a winning margin of 14 points, our sorted list is now

8, 14, **31**, **32**, 56, 56

and there are *two* middle numbers, 31 and 32. The median is defined as the average of those two numbers, which is of course 31.5. As before, it's very tedious to do this by hand when you've got lots of numbers. To illustrate this, here's what happens when you use R to sort all 176 winning margins. First, I'll use the `sort()` function (discussed in Chapter 7) to display the winning margins in increasing numerical order:

```
> sort( x = afl.margins )
 [1]  0   0   1   1   1   1   1   2   2   2   3   3   3   3   3   3
[14]  3   3   3   4   4   4   5   6   7   7   7   8   8   8   8   8
[27]  8   9   9   9   9   9   9   10  10  10  10  10  10  10  11
[40] 11  11  12  12  12  13  14  14  14  15  15  16  16  16  16
[53] 18  19  19  19  19  19  19  20  20  20  20  21  21  21  22
[66] 22  23  23  23  24  24  25  25  25  26  26  26  26  26  27
[79] 27  28  28  29  29  29  29  29  29  29  29  30  31  32  32
[92] 33  35  35  35  35  35  36  36  36  36  36  36  36  37  38
[105] 38  38  38  38  38  39  39  40  41  42  42  43  43  44  44
[118] 44  44  44  47  47  47  47  48  48  48  49  49  49  50  50
[131] 50  50  52  52  53  53  54  54  55  55  55  55  56  56
[144] 56  57  60  61  61  63  64  65  65  66  67  68  70
[157] 71  71  72  73  75  75  76  81  82  82  83  84  89
[170] 94  95  98 101 104 108 116
```

The middle values (which I've shown in bold) are 30 and 31, so the median winning margin for 2010 was 30.5 points. In real life, of course, no-one actually calculates the median by sorting the data and then looking for the middle value. In real life, we use the `median` command:

```
> median( x = afl.margins )
[1] 30.5
```

which outputs the median value of 30.5.

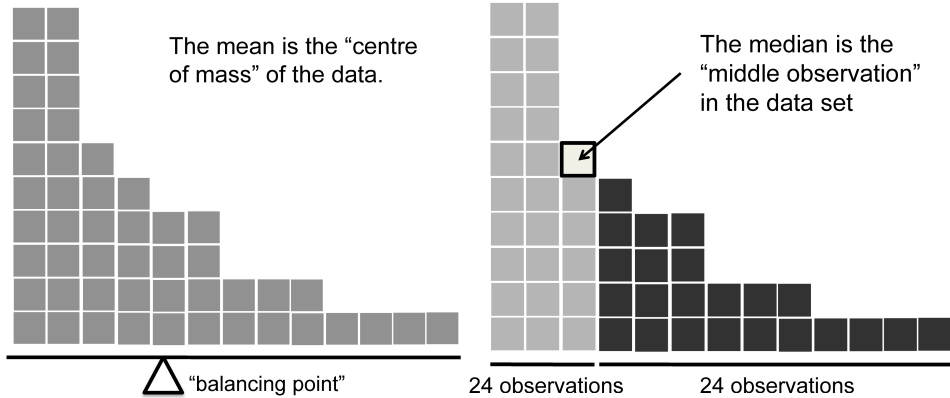


Figure 5.2: An illustration of the difference between how the mean and the median should be interpreted. The mean is basically the “centre of gravity” of the data set: if you imagine that the histogram of the data is a solid object, then the point on which you could balance it (as if on a see-saw) is the mean. In contrast, the median is the “middle value” in the data. What this implies, as far as which one you should use, depends a little on what type of data you’ve got and what you’re trying to achieve. As a rough guide:

- If your data are nominal scale, you probably shouldn’t be using either the mean or the median. Both the mean and the median rely on the idea that the numbers assigned to values are meaningful. If the numbering scheme is arbitrary, then it’s probably best to use the mode (Section 5.1.7) instead.
- If your data are ordinal scale, you’re more likely to want to use the median than the mean. The median only makes use of the order information in your data (i.e., which numbers are bigger), but doesn’t depend on the precise numbers involved. That’s exactly the situation that applies when your data are ordinal scale. The mean, on the other hand, makes use of the precise numeric values assigned to the observations, so it’s not really appropriate for ordinal data.
- For interval and ratio scale data, either one is generally acceptable. Which one you pick depends a bit on what you’re trying to achieve. The mean has the advantage that it uses all the information in the data (which is useful when you don’t have a lot of data), but it’s very sensitive to extreme values, as we’ll see in Section 5.1.6.

Let’s expand on that last part a little. One consequence is that there’s systematic differences between the mean and the median when the histogram is asymmetric (skewed; see Section 5.3). This is illustrated in Figure 5.2: notice that the median (right hand side) is located closer to the “body” of the histogram, whereas the mean (left hand side) gets dragged towards the “tail” (where the extreme values are). To give a concrete example, suppose Bob (income \$50,000), Kate (income \$60,000) and Jane (income \$65,000) are sitting at a table: the average income at the table is \$58,333 and the median income is \$60,000. Then Bill sits down with them (income \$100,000,000). The average income has now jumped to \$25,043,750 but

the median rises only to \$62,500. If you're interested in looking at the overall income at the table, the mean might be the right answer; but if you're interested in what counts as a typical income at the table, the median would be a better choice here.

5.1.5 A real life example

To try to get a sense of why you need to pay attention to the differences between the mean and the median, let's consider a real life example. Since I tend to mock journalists for their poor scientific and statistical knowledge, I should give credit where credit is due. This is from an excellent article on the ABC news website⁴ 24 September, 2010:

Senior Commonwealth Bank executives have travelled the world in the past couple of weeks with a presentation showing how Australian house prices, and the key price to income ratios, compare favourably with similar countries. "Housing affordability has actually been going sideways for the last five to six years," said Craig James, the chief economist of the bank's trading arm, CommSec.

This probably comes as a huge surprise to anyone with a mortgage, or who wants a mortgage, or pays rent, or isn't completely oblivious to what's been going on in the Australian housing market over the last several years. Back to the article:

CBA has waged its war against what it believes are housing doomsayers with graphs, numbers and international comparisons. In its presentation, the bank rejects arguments that Australia's housing is relatively expensive compared to incomes. It says Australia's house price to household income ratio of 5.6 in the major cities, and 4.3 nationwide, is comparable to many other developed nations. It says San Francisco and New York have ratios of 7, Auckland's is 6.7, and Vancouver comes in at 9.3.

More excellent news! Except, the article goes on to make the observation that...

Many analysts say that has led the bank to use misleading figures and comparisons. If you go to page four of CBA's presentation and read the source information at the bottom of the graph and table, you would notice there is an additional source on the international comparison – Demographia. However, if the Commonwealth Bank had also used Demographia's analysis of Australia's house price to income ratio, it would have come up with a figure closer to 9 rather than 5.6 or 4.3

That's, um, a rather serious discrepancy. One group of people say 9, another says 4-5. Should we just split the difference, and say the truth lies somewhere in between? Absolutely not: this is a situation where there is a right answer and a wrong answer. Demographia are correct, and the Commonwealth Bank is incorrect. As the article points out

[An] obvious problem with the Commonwealth Bank's domestic price to income figures is they compare average incomes with median house prices (unlike the Demographia figures that compare median incomes to median prices). The median is the mid-point, effectively cutting out the highs and lows, and that means the average is generally higher when it comes to incomes and asset prices, because it includes the earnings of Australia's wealthiest people. To put it another way: the Commonwealth Bank's figures count Ralph Norris' multi-million dollar pay packet on the income side, but not his (no doubt) very expensive house in the property price figures, thus understating the house price to income ratio for middle-income Australians.

⁴www.abc.net.au/news/stories/2010/09/24/3021480.htm

Couldn't have put it better myself. The way that Demographia calculated the ratio is the right thing to do. The way that the Bank did it is incorrect. As for why an extremely quantitatively sophisticated organisation such as a major bank made such an elementary mistake, well... I can't say for sure, since I have no special insight into their thinking, but the article itself does happen to mention the following facts, which may or may not be relevant:

[As] Australia's largest home lender, the Commonwealth Bank has one of the biggest vested interests in house prices rising. It effectively owns a massive swathe of Australian housing as security for its home loans as well as many small business loans.

My, my.

5.1.6 Trimmed mean

One of the fundamental rules of applied statistics is that the data are messy. Real life is never simple, and so the data sets that you obtain are never as straightforward as the statistical theory says.⁵ This can have awkward consequences. To illustrate, consider this rather strange looking data set:

$$-100, 2, 3, 4, 5, 6, 7, 8, 9, 10$$

If you were to observe this in a real life data set, you'd probably suspect that something funny was going on with the -100 value. It's probably an **outlier**, a value that doesn't really belong with the others. You might consider removing it from the data set entirely, and in this particular case I'd probably agree with that course of action. In real life, however, you don't always get such cut-and-dried examples. For instance, you might get this instead:

$$-15, 2, 3, 4, 5, 6, 7, 8, 9, 12$$

The -15 looks a bit suspicious, but not anywhere near as much as that -100 did. In this case, it's a little trickier. It *might* be a legitimate observation, it *might* not.

When faced with a situation where some of the most extreme-valued observations might not be quite trustworthy, the mean is not necessarily a good measure of central tendency. It is highly sensitive to one or two extreme values, and is thus not considered to be a **robust** measure. One remedy that we've seen is to use the median. A more general solution is to use a "trimmed mean". To calculate a trimmed mean, what you do is "discard" the most extreme examples on both ends (i.e., the largest and the smallest), and then take the mean of everything else. The goal is to preserve the best characteristics of the mean and the median: just like a median, you aren't highly influenced by extreme outliers, but like the mean, you "use" more than one of the observations. Generally, we describe a trimmed mean in terms of the percentage of observation on either side that are discarded. So, for instance, a 10% trimmed mean discards the largest 10% of the observations *and* the smallest 10% of the observations, and then takes the mean of the remaining 80% of the observations. Not surprisingly, the 0% trimmed mean is just the regular mean, and the 50% trimmed mean is the median. In that sense, trimmed means provide a whole family of central tendency measures that span the range from the mean to the median.

For our toy example above, we have 10 observations, and so a 10% trimmed mean is calculated by ignoring the largest value (i.e., 12) and the smallest value (i.e., -15) and taking the mean of the remaining values. First, let's enter the data

```
> dataset <- c( -15,2,3,4,5,6,7,8,9,12 )
```

⁵Or at least, the basic statistical theory – these days there is a whole subfield of statistics called *robust statistics* that tries to grapple with the messiness of real data and develop theory that can cope with it.

Next, let's calculate means and medians:

```
> mean( x = dataset )
[1] 4.1
> median( x = dataset )
[1] 5.5
```

That's a fairly substantial difference, but I'm tempted to think that the mean is being influenced a bit too much by the extreme values at either end of the data set, especially the -15 one. So let's just try trimming the mean a bit. If I take a 10% trimmed mean, we'll drop the extreme values on either side, and take the mean of the rest:

```
> mean( x = dataset, trim = .1)
[1] 5.5
```

which in this case gives exactly the same answer as the median. Note that, to get a 10% trimmed mean you write `trim = .1`, not `trim = 10`. In any case, let's finish up by calculating the 5% trimmed mean for the `afl.margins` data,

```
> mean( x = afl.margins, trim = .05)
[1] 33.75
```

5.1.7 Mode

The mode of a sample is very simple: it is the value that occurs most frequently. To illustrate the mode using the AFL data, let's examine a different aspect to the data set. Who has played in the most finals? The `afl.finalists` variable is a factor that contains the name of every team that played in any AFL final from 1987-2010, so let's have a look at it:

```
> print( afl.finalists )
 [1] Hawthorn      Melbourne     Carlton
 [4] Melbourne     Hawthorn     Carlton
 [7] Melbourne     Carlton     Hawthorn
 [10] Melbourne    Melbourne    Hawthorn

 BLAH BLAH BLAH

 [391] St Kilda    Hawthorn     Western Bulldogs
 [394] Carlton     Fremantle   Sydney
 [397] Geelong     Western Bulldogs St Kilda
 [400] St Kilda    Adelaide    Brisbane Carlton ... Western Bulldogs
 17 Levels: Adelaide Brisbane Carlton ... Western Bulldogs
```

What we *could* do is read through all 400 entries, and count the number of occasions on which each team name appears in our list of finalists, thereby producing a **frequency table**. However, that would be mindless and boring: exactly the sort of task that computers are great at. So let's use the `table()` function (discussed in more detail in Section 7.1) to do this task for us:

```
> table( afl.finalists )
afl.finalists
Adelaide      Brisbane      Carlton      Collingwood
 26           25            26           28
```

Essendon	Fitzroy	Fremantle	Geelong
32	0	6	39
Hawthorn	Melbourne	North Melbourne	Port Adelaide
27	28	28	17
Richmond	St Kilda	Sydney	West Coast
6	24	26	38
Western Bulldogs			
24			

Now that we have our frequency table, we can just look at it and see that, over the 24 years for which we have data, Geelong has played in more finals than any other team. Thus, the mode of the `finalists` data is "Geelong". The core packages in R don't have a function for calculating the mode⁶. However, I've included a function in the `lsr` package that does this. The function is called `modeOf()`, and here's how you use it:

```
> modeOf( x = afl.finalists )
[1] "Geelong"
```

There's also a function called `maxFreq()` that tells you what the modal frequency is. If we apply this function to our `finalists` data, we obtain the following:

```
> maxFreq( x = afl.finalists )
[1] 39
```

Taken together, we observe that Geelong (39 finals) played in more finals than any other team during the 1987-2010 period.

One last point to make with respect to the mode. While it's generally true that the mode is most often calculated when you have nominal scale data (because means and medians are useless for those sorts of variables), there are some situations in which you really do want to know the mode of an ordinal, interval or ratio scale variable. For instance, let's go back to thinking about our `afl.margins` variable. This variable is clearly ratio scale (if it's not clear to you, it may help to re-read Section 2.2), and so in most situations the mean or the median is the measure of central tendency that you want. But consider this scenario... a friend of yours is offering a bet. They pick a football game at random, and (without knowing who is playing) you have to guess the *exact* margin. If you guess correctly, you win \$50. If you don't, you lose \$1. There are no consolation prizes for "almost" getting the right answer. You have to guess exactly the right margin⁷ For this bet, the mean and the median are completely useless to you. It is the mode that you should bet on. So, we calculate this modal value

```
> modeOf( x = afl.margins )
[1] 3
> maxFreq( x = afl.margins )
[1] 8
```

So the 2010 data suggest you should bet on a 3 point margin, and since this was observed in 8 of the 176 game (4.5% of games) the odds are firmly in your favour.

⁶As we saw earlier, it *does* have a function called `mode()`, but it does something completely different.

⁷This is called a "0-1 loss function", meaning that you either win (1) or you lose (0), with no middle ground.

Measures of variability

The statistics that we've discussed so far all relate to *central tendency*. That is, they all talk about which values are “in the middle” or “popular” in the data. However, central tendency is not the only type of summary statistic that we want to calculate. The second thing that we really want is a measure of the **variability** of the data. That is, how “spread out” are the data? How “far” away from the mean or median do the observed values tend to be? For now, let's assume that the data are interval or ratio scale, so we'll continue to use the `afl.margins` data. We'll use this data to discuss several different measures of spread, each with different strengths and weaknesses.

5.2.1 Range

The **range** of a variable is very simple: it's the biggest value minus the smallest value. For the AFL winning margins data, the maximum value is 116, and the minimum value is 0. We can calculate these values in R using the `max()` and `min()` functions:

```
> max( afl.margins )
> min( afl.margins )
```

where I've omitted the output because it's not interesting. The other possibility is to use the `range()` function; which outputs both the minimum value and the maximum value in a vector, like this:

```
> range( afl.margins )
[1] 0 116
```

Although the range is the simplest way to quantify the notion of “variability”, it's one of the worst. Recall from our discussion of the mean that we want our summary measure to be robust. If the data set has one or two extremely bad values in it, we'd like our statistics not to be unduly influenced by these cases. If we look once again at our toy example of a data set containing very extreme outliers...

-100, 2, 3, 4, 5, 6, 7, 8, 9, 10

... it is clear that the range is not robust, since this has a range of 110, but if the outlier were removed we would have a range of only 8.

5.2.2 Interquartile range

The **interquartile range** (IQR) is like the range, but instead of calculating the difference between the biggest and smallest value, it calculates the difference between the 25th quantile and the 75th quantile. Probably you already know what a **quantile** is (they're more commonly called percentiles), but if not: the 10th percentile of a data set is the smallest number x such that 10% of the data is less than x . In fact, we've already come across the idea: the median of a data set is its 50th quantile / percentile! R actually provides you with a way of calculating quantiles, using the (surprise, surprise) `quantile()` function. Let's use it to calculate the median AFL winning margin:

```
> quantile( x = afl.margins, probs = .5)
50%
30.5
```

And not surprisingly, this agrees with the answer that we saw earlier with the `median()` function. Now, we can actually input lots of quantiles at once, by specifying a vector for the `probs` argument. So let's do that, and get the 25th and 75th percentile:

```
> quantile( x = afl.margins, probs = c(.25, .75) )
25%    75%
12.75 50.50
```

And, by noting that $50.5 - 12.75 = 37.75$, we can see that the interquartile range for the 2010 AFL winning margins data is 37.75. Of course, that seems like too much work to do all that typing, so R has a built in function called `IQR()` that we can use:

```
> IQR( x = afl.margins )
[1] 37.75
```

While it's obvious how to interpret the range, it's a little less obvious how to interpret the IQR. The simplest way to think about it is like this: the interquartile range is the range spanned by the "middle half" of the data. That is, one quarter of the data falls below the 25th percentile, one quarter of the data is above the 75th percentile, leaving the "middle half" of the data lying in between the two. And the IQR is the range covered by that middle half.

5.2.3 Mean absolute deviation

The two measures we've looked at so far, the range and the interquartile range, both rely on the idea that we can measure the spread of the data by looking at the quantiles of the data. However, this isn't the only way to think about the problem. A different approach is to select a meaningful reference point (usually the mean or the median) and then report the "typical" deviations from that reference point. What do we mean by "typical" deviation? Usually, the mean or median value of these deviations! In practice, this leads to two different measures, the "mean absolute deviation (from the mean)" and the "median absolute deviation (from the median)". From what I've read, the measure based on the median seems to be used in statistics, and does seem to be the better of the two, but to be honest I don't think I've seen it used much in psychology. The measure based on the mean does occasionally show up in psychology though. In this section I'll talk about the first one, and I'll come back to talk about the second one later.

Since the previous paragraph might sound a little abstract, let's go through the **mean absolute deviation** from the mean a little more slowly. One useful thing about this measure is that the name actually tells you exactly how to calculate it. Let's think about our AFL winning margins data, and once again we'll start by pretending that there's only 5 games in total, with winning margins of 56, 31, 56, 8 and 32. Since our calculations rely on an examination of the deviation from some reference point (in this case the mean), the first thing we need to calculate is the mean, \bar{X} . For these five observations, our mean is $\bar{X} = 36.6$. The next step is to convert each of our observations X_i into a deviation score. We do this by calculating the difference between the observation X_i and the mean \bar{X} . That is, the deviation score is defined to be $X_i - \bar{X}$. For the first observation in our sample, this is equal to $56 - 36.6 = 19.4$. Okay, that's simple enough. The next step in the process is to convert these deviations to absolute deviations. As we discussed earlier when talking about the `abs()` function in R (Section 3.5), we do this by converting any negative values to positive ones. Mathematically, we would denote the absolute value of -3 as $|-3|$, and so we say that $|-3| = 3$. We use the absolute value function here because we don't really care whether the value is higher than the mean or lower than the mean, we're just interested in how *close* it is to the mean. To help make this process as obvious as possible, the table below shows these calculations for all five observations:

English: notation:	which game <i>i</i>	value X_i	deviation from mean $X_i - \bar{X}$	absolute deviation $ X_i - \bar{X} $
	1	56	19.4	19.4
	2	31	-5.6	5.6
	3	56	19.4	19.4
	4	8	-28.6	28.6
	5	32	-4.6	4.6

Now that we have calculated the absolute deviation score for every observation in the data set, all that we have to do is calculate the mean of these scores. Let's do that:

$$\frac{19.4 + 5.6 + 19.4 + 28.6 + 4.6}{5} = 15.52$$

And we're done. The mean absolute deviation for these five scores is 15.52.

However, while our calculations for this little example are at an end, we do have a couple of things left to talk about. Firstly, we should really try to write down a proper mathematical formula. But in order to do this I need some mathematical notation to refer to the mean absolute deviation. Irritatingly, "mean absolute deviation" and "median absolute deviation" have the same acronym (MAD), which leads to a certain amount of ambiguity, and since R tends to use MAD to refer to the median absolute deviation, I'd better come up with something different for the mean absolute deviation. Sigh. What I'll do is use AAD instead, short for *average* absolute deviation. Now that we have some unambiguous notation, here's the formula that describes what we just calculated:

$$AAD(X) = \frac{1}{N} \sum_{i=1}^N |X_i - \bar{X}|$$

The last thing we need to talk about is how to calculate AAD in R. One possibility would be to do everything using low level commands, laboriously following the same steps that I used when describing the calculations above. However, that's pretty tedious. You'd end up with a series of commands that might look like this:

```
> X <- c(56, 31, 56, 8, 32)      # enter the data
> X.bar <- mean(X)                # step 1. the mean of the data
> AD <- abs(X - X.bar)            # step 2. the absolute deviations from the mean
> AAD <- mean(AD)                 # step 3. the mean absolute deviations
> print(AAD)                      # print the results
[1] 15.52
```

Each of those commands is pretty simple, but there's just too many of them. And because I find that to be too much typing, the `lsr` package has a very simple function called `aad()` that does the calculations for you. If we apply the `aad()` function to our data, we get this:

```
> aad(X)
[1] 15.52
```

No surprises there.

5.2.4 Variance

Although the mean absolute deviation measure has its uses, it's not the best measure of variability to use. From a purely mathematical perspective, there are some solid reasons to prefer squared deviations

rather than absolute deviations. If we do that, we obtain a measure is called the **variance**, which has a lot of really nice statistical properties that I'm going to ignore,⁸ and one massive psychological flaw that I'm going to make a big deal out of in a moment. The variance of a data set X is sometimes written as $\text{Var}(X)$, but it's more commonly denoted s^2 (the reason for this will become clearer shortly). The formula that we use to calculate the variance of a set of observations is as follows:

$$\text{Var}(X) = \frac{1}{N} \sum_{i=1}^N (X_i - \bar{X})^2$$

As you can see, it's basically the same formula that we used to calculate the mean absolute deviation, except that instead of using “absolute deviations” we use “squared deviations”. It is for this reason that the variance is sometimes referred to as the “mean square deviation”.

Now that we've got the basic idea, let's have a look at a concrete example. Once again, let's use the first five AFL games as our data. If we follow the same approach that we took last time, we end up with the following table:

English: maths:	which game i	value X_i	deviation from mean $X_i - \bar{X}$	squared deviation $(X_i - \bar{X})^2$
	1	56	19.4	376.36
	2	31	-5.6	31.36
	3	56	19.4	376.36
	4	8	-28.6	817.96
	5	32	-4.6	21.16

That last column contains all of our squared deviations, so all we have to do is average them. If we do that by typing all the numbers into R by hand...

```
> ( 376.36 + 31.36 + 376.36 + 817.96 + 21.16 ) / 5
[1] 324.64
```

... we end up with a variance of 324.64. Exciting, isn't it? For the moment, let's ignore the burning question that you're all probably thinking (i.e., what the heck does a variance of 324.64 actually mean?) and instead talk a bit more about how to do the calculations in R, because this will reveal something very weird.

As always, we want to avoid having to type in a whole lot of numbers ourselves. And as it happens, we have the vector `X` lying around, which we created in the previous section. With this in mind, we can calculate the variance of `X` by using the following command,

```
> mean( (X - mean(X))^2 )
[1] 324.64
```

and as usual we get the same answer as the one that we got when we did everything by hand. However, I *still* think that this is too much typing. Fortunately, R has a built in function called `var()` which does calculate variances. So we could also do this...

```
> var(X)
[1] 405.8
```

⁸Well, I will very briefly mention the one that I think is coolest, for a very particular definition of “cool”, that is. Variances are *additive*. Here's what that means: suppose I have two variables X and Y , whose variances are $\text{Var}(X)$ and $\text{Var}(Y)$ respectively. Now imagine I want to define a new variable Z that is the sum of the two, $Z = X + Y$. As it turns out, the variance of Z is equal to $\text{Var}(X) + \text{Var}(Y)$. This is a *very* useful property, but it's not true of the other measures that I talk about in this section.

and you get the same... no, wait... you get a completely *different* answer. That's just weird. Is R broken? Is this a typo? Is Dan an idiot?

As it happens, the answer is no.⁹ It's not a typo, and R is not making a mistake. To get a feel for what's happening, let's stop using the tiny data set containing only 5 data points, and switch to the full set of 176 games that we've got stored in our `afl.margins` vector. First, let's calculate the variance by using the formula that I described above:

```
> mean( (afl.margins - mean(afl.margins) )^2)
[1] 675.9718
```

Now let's use the `var()` function:

```
> var( afl.margins )
[1] 679.8345
```

Hm. These two numbers are very similar this time. That seems like too much of a coincidence to be a mistake. And of course it isn't a mistake. In fact, it's very simple to explain what R is doing here, but slightly trickier to explain *why* R is doing it. So let's start with the "what". What R is doing is evaluating a slightly different formula to the one I showed you above. Instead of averaging the squared deviations, which requires you to divide by the number of data points N , R has chosen to divide by $N - 1$. In other words, the formula that R is using is this one

$$\frac{1}{N-1} \sum_{i=1}^N (X_i - \bar{X})^2$$

It's easy enough to verify that this is what's happening, as the following command illustrates:

```
> sum( (X-mean(X))^2 ) / 4
[1] 405.8
```

This is the same answer that R gave us originally when we calculated `var(X)` originally. So that's the *what*. The real question is *why* R is dividing by $N - 1$ and not by N . After all, the variance is supposed to be the *mean* squared deviation, right? So shouldn't we be dividing by N , the actual number of observations in the sample? Well, yes, we should. However, as we'll discuss in Chapter 10, there's a subtle distinction between "describing a sample" and "making guesses about the population from which the sample came". Up to this point, it's been a distinction without a difference. Regardless of whether you're describing a sample or drawing inferences about the population, the mean is calculated exactly the same way. Not so for the variance, or the standard deviation, or for many other measures besides. What I outlined to you initially (i.e., take the actual average, and thus divide by N) assumes that you literally intend to calculate the variance of the sample. Most of the time, however, you're not terribly interested in the sample *in and of itself*. Rather, the sample exists to tell you something about the world. If so, you're actually starting to move away from calculating a "sample statistic", and towards the idea of estimating a "population parameter". However, I'm getting ahead of myself. For now, let's just take it on faith that R knows what it's doing, and we'll revisit the question later on when we talk about estimation in Chapter 10.

Okay, one last thing. This section so far has read a bit like a mystery novel. I've shown you how to calculate the variance, described the weird " $N - 1$ " thing that R does and hinted at the reason why it's there, but I haven't mentioned the single most important thing... how do you *interpret* the variance? Descriptive statistics are supposed to describe things, after all, and right now the variance is really just a gibberish number. Unfortunately, the reason why I haven't given you the human-friendly interpretation of

⁹With the possible exception of the third question.

the variance is that there really isn't one. This is the most serious problem with the variance. Although it has some elegant mathematical properties that suggest that it really is a fundamental quantity for expressing variation, it's completely useless if you want to communicate with an actual human... variances are completely uninterpretable in terms of the original variable! All the numbers have been squared, and they don't mean anything anymore. This is a huge issue. For instance, according to the table I presented earlier, the margin in game 1 was "376.36 points-squared higher than the average margin". This is *exactly* as stupid as it sounds; and so when we calculate a variance of 324.64, we're in the same situation. I've watched a lot of footy games, and never has anyone referred to "points squared". It's *not* a real unit of measurement, and since the variance is expressed in terms of this gibberish unit, it is totally meaningless to a human.

5.2.5 Standard deviation

Okay, suppose that you like the idea of using the variance because of those nice mathematical properties that I haven't talked about, but – since you're a human and not a robot – you'd like to have a measure that is expressed in the same units as the data itself (i.e., points, not points-squared). What should you do? The solution to the problem is obvious: take the square root of the variance, known as the **standard deviation**, also called the "root mean squared deviation", or RMSD. This solves our problem fairly neatly: while nobody has a clue what "a variance of 324.68 points-squared" really means, it's much easier to understand "a standard deviation of 18.01 points", since it's expressed in the original units. It is traditional to refer to the standard deviation of a sample of data as s , though "sd" and "std dev." are also used at times. Because the standard deviation is equal to the square root of the variance, you probably won't be surprised to see that the formula is:

$$s = \sqrt{\frac{1}{N} \sum_{i=1}^N (X_i - \bar{X})^2}$$

and the R function that we use to calculate it is `sd()`. However, as you might have guessed from our discussion of the variance, what R actually calculates is slightly different to the formula given above. Just like we saw with the variance, what R calculates is a version that divides by $N - 1$ rather than N . For reasons that will make sense when we return to this topic in Chapter 10 I'll refer to this new quantity as $\hat{\sigma}$ (read as: "sigma hat"), and the formula for this is

$$\hat{\sigma} = \sqrt{\frac{1}{N-1} \sum_{i=1}^N (X_i - \bar{X})^2}$$

With that in mind, calculating standard deviations in R is simple:

```
> sd(afl.margins)
[1] 26.07364
```

Interpreting standard deviations is slightly more complex. Because the standard deviation is derived from the variance, and the variance is a quantity that has little to no meaning that makes sense to us humans, the standard deviation doesn't have a simple interpretation. As a consequence, most of us just rely on a simple rule of thumb: in general, you should expect 68% of the data to fall within 1 standard deviation of the mean, 95% of the data to fall within 2 standard deviation of the mean, and 99.7% of the data to fall within 3 standard deviations of the mean. This rule tends to work pretty well most of the time, but it's not exact: it's actually calculated based on an *assumption* that the histogram is symmetric and "bell shaped".¹⁰ As you can tell from looking at the AFL winning margins histogram in

¹⁰Strictly, the assumption is that the data are *normally* distributed, which is an important concept that we'll discuss more in Chapter 9, and will turn up over and over again later in the book.

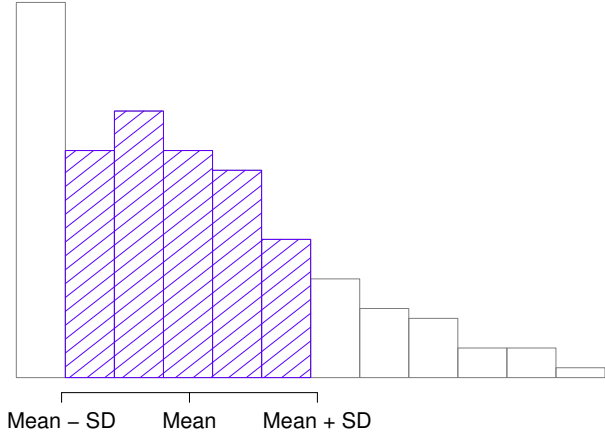


Figure 5.3: An illustration of the standard deviation, applied to the AFL winning margins data. The shaded bars in the histogram show how much of the data fall within one standard deviation of the mean. In this case, 65.3% of the data set lies within this range, which is pretty consistent with the “approximately 68% rule” discussed in the main text.

Figure 5.1, this isn’t exactly true of our data! Even so, the rule is approximately correct. As it turns out, 65.3% of the AFL margins data fall within one standard deviation of the mean. This is shown visually in Figure 5.3.

5.2.6 Median absolute deviation

The last measure of variability that I want to talk about is the **median absolute deviation** (MAD). The basic idea behind MAD is very simple, and is pretty much identical to the idea behind the mean absolute deviation (Section 5.2.3). The difference is that you use the median everywhere. If we were to frame this idea as a pair of R commands, they would look like this:

```
> # mean absolute deviation from the mean:
> mean( abs(afl.margins - mean(afl.margins)) )
[1] 21.10124

> # *median* absolute deviation from the *median*:
> median( abs(afl.margins - median(afl.margins)) )
[1] 19.5
```

This has a straightforward interpretation: every observation in the data set lies some distance away from the typical value (the median). So the MAD is an attempt to describe a *typical deviation from a typical value* in the data set. It wouldn’t be unreasonable to interpret the MAD value of 19.5 for our AFL data by saying something like this:

The median winning margin in 2010 was 30.5, indicating that a typical game involved a

winning margin of about 30 points. However, there was a fair amount of variation from game to game: the MAD value was 19.5, indicating that a typical winning margin would differ from this median value by about 19-20 points.

As you'd expect, R has a built in function for calculating MAD, and you will be shocked no doubt to hear that it's called `mad()`. However, it's a little bit more complicated than the functions that we've been using previously. If you want to use it to calculate MAD in the exact same way that I have described it above, the command that you need to use specifies two arguments: the data set itself `x`, and a `constant` that I'll explain in a moment. For our purposes, the constant is 1, so our command becomes

```
> mad( x = afl.margins, constant = 1 )
[1] 19.5
```

Apart from the weirdness of having to type that `constant = 1` part, this is pretty straightforward.

Okay, so what exactly is this `constant = 1` argument? I won't go into all the details here, but here's the gist. Although the "raw" MAD value that I've described above is completely interpretable on its own terms, that's not actually how it's used in a lot of real world contexts. Instead, what happens a lot is that the researcher *actually* wants to calculate the standard deviation. However, in the same way that the mean is very sensitive to extreme values, the standard deviation is vulnerable to the exact same issue. So, in much the same way that people sometimes use the median as a "robust" way of calculating "something that is like the mean", it's not uncommon to use MAD as a method for calculating "something that is like the standard deviation". Unfortunately, the *raw* MAD value doesn't do this. Our raw MAD value is 19.5, and our standard deviation was 26.07. However, what some clever person has shown is that, under certain assumptions¹¹, you can multiply the raw MAD value by 1.4826 and obtain a number that is directly comparable to the standard deviation. As a consequence, the default value of `constant` is 1.4826, and so when you use the `mad()` command without manually setting a value, here's what you get:

```
> mad( afl.margins )
[1] 28.9107
```

I should point out, though, that if you want to use this "corrected" MAD value as a robust version of the standard deviation, you really are relying on the assumption that the data are (or at least, are "supposed to be" in some sense) symmetric and basically shaped like a bell curve. That's really *not* true for our `afl.margins` data, so in this case I wouldn't try to use the MAD value this way.

5.2.7 Which measure to use?

We've discussed quite a few measures of spread (range, IQR, MAD, variance and standard deviation), and hinted at their strengths and weaknesses. Here's a quick summary:

- *Range*. Gives you the full spread of the data. It's very vulnerable to outliers, and as a consequence it isn't often used unless you have good reasons to care about the extremes in the data.
- *Interquartile range*. Tells you where the "middle half" of the data sits. It's pretty robust, and complements the median nicely. This is used a lot.
- *Mean absolute deviation*. Tells you how far "on average" the observations are from the mean. It's very interpretable, but has a few minor issues (not discussed here) that make it less attractive to statisticians than the standard deviation. Used sometimes, but not often.

¹¹The assumption again being that the data are normally-distributed!

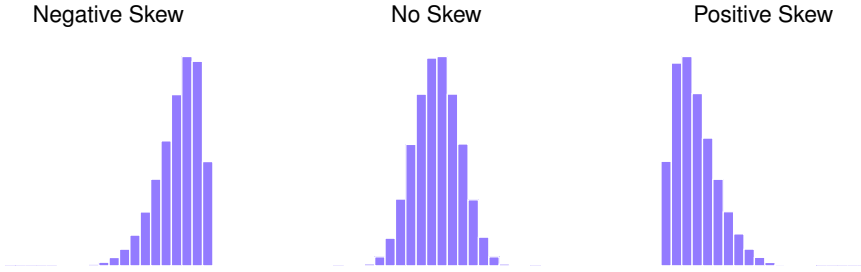


Figure 5.4: An illustration of skewness. On the left we have a negatively skewed data set (skewness = $-.93$), in the middle we have a data set with no skew (technically, skewness = $-.006$), and on the right we have a positively skewed data set (skewness = $.93$).

- *Variance.* Tells you the average squared deviation from the mean. It's mathematically elegant, and is probably the “right” way to describe variation around the mean, but it's completely uninterpretable because it doesn't use the same units as the data. Almost never used except as a mathematical tool; but it's buried “under the hood” of a very large number of statistical tools.
- *Standard deviation.* This is the square root of the variance. It's fairly elegant mathematically, and it's expressed in the same units as the data so it can be interpreted pretty well. In situations where the mean is the measure of central tendency, this is the default. This is by far the most popular measure of variation.
- *Median absolute deviation.* The typical (i.e., median) deviation from the median value. In the raw form it's simple and interpretable; in the corrected form it's a robust way to estimate the standard deviation, for some kinds of data sets. Not used very often, but it does get reported sometimes.

In short, the IQR and the standard deviation are easily the two most common measures used to report the variability of the data; but there are situations in which the others are used. I've described all of them in this book because there's a fair chance you'll run into most of these somewhere.

5.3 _____

Skew and kurtosis

There are two more descriptive statistics that you will sometimes see reported in the psychological literature, known as skew and kurtosis. In practice, neither one is used anywhere near as frequently as the measures of central tendency and variability that we've been talking about. Skew is pretty important, so you do see it mentioned a fair bit; but I've actually never seen kurtosis reported in a scientific article to date.

Since it's the more interesting of the two, let's start by talking about the **skewness**. Skewness is basically a measure of asymmetry, and the easiest way to explain it is by drawing some pictures. As Figure 5.4 illustrates, if the data tend to have a lot of extreme small values (i.e., the lower tail is “longer” than the upper tail) and not so many extremely large values (left panel), then we say that the data are

negatively skewed. On the other hand, if there are more extremely large values than extremely small ones (right panel) we say that the data are *positively skewed*. That's the qualitative idea behind skewness. The actual formula for the skewness of a data set is as follows

$$\text{skewness}(X) = \frac{1}{N\hat{\sigma}^3} \sum_{i=1}^N (X_i - \bar{X})^3$$

where N is the number of observations, \bar{X} is the sample mean, and $\hat{\sigma}$ is the standard deviation (the “divide by $N - 1$ ” version, that is). Perhaps more helpfully, it might be useful to point out that the `psych` package contains a `skew()` function that you can use to calculate skewness. So if we wanted to use this function to calculate the skewness of the `afl.margins` data, we'd first need to load the package

```
> library(psych)
```

which now makes it possible to use the following command:

```
> skew(x = afl.margins)
[1] 0.7671555
```

Not surprisingly, it turns out that the AFL winning margins data is fairly skewed.

The final measure that is sometimes referred to, though very rarely in practice, is the **kurtosis** of a data set. Put simply, kurtosis is a measure of the “pointiness” of a data set, as illustrated in Figure 5.5. By convention, we say that the “normal curve” (black lines) has zero kurtosis, so the pointiness of a data set is assessed relative to this curve. In this Figure, the data on the left are not pointy enough, so the kurtosis is negative and we call the data *platykurtic*. The data on the right are too pointy, so the kurtosis is positive and we say that the data is *leptokurtic*. But the data in the middle are just pointy enough, so we say that it is *mesokurtic* and has kurtosis zero. This is summarised in the table below:

informal term	technical name	kurtosis value
“too flat”	platykurtic	negative
“just pointy enough”	mesokurtic	zero
“too pointy”	leptokurtic	positive

The equation for kurtosis is pretty similar in spirit to the formulas we've seen already for the variance and the skewness; except that where the variance involved squared deviations and the skewness involved cubed deviations, the kurtosis involves raising the deviations to the fourth power:¹²

$$\text{kurtosis}(X) = \frac{1}{N\hat{\sigma}^4} \sum_{i=1}^N (X_i - \bar{X})^4 - 3$$

I know, it's not terribly interesting to me either. More to the point, the `psych` package has a function called `kurtosi()` that you can use to calculate the kurtosis of your data. For instance, if we were to do this for the AFL margins,

```
> kurtosi(x = afl.margins)
[1] 0.02962633
```

we discover that the AFL winning margins data are just pointy enough.

¹²The “-3” part is something that statisticians tack on to ensure that the normal curve has kurtosis zero. It looks a bit stupid, just sticking a “-3” at the end of the formula, but there are good mathematical reasons for doing this.

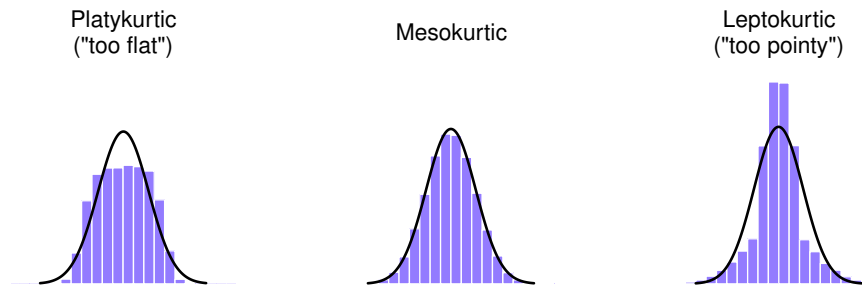


Figure 5.5: An illustration of kurtosis. On the left, we have a “platykurtic” data set (kurtosis = -0.95), meaning that the data set is “too flat”. In the middle we have a “mesokurtic” data set (kurtosis is almost exactly 0), which means that the pointiness of the data is just about right. Finally, on the right, we have a “leptokurtic” data set (kurtosis = 2.12) indicating that the data set is “too pointy”. Note that kurtosis is measured with respect to a normal curve (black line).

.....

5.4

Getting an overall summary of a variable

Up to this point in the chapter I’ve explained several different summary statistics that are commonly used when analysing data, along with specific functions that you can use in R to calculate each one. However, it’s kind of annoying to have to separately calculate means, medians, standard deviations, skewness etc. Wouldn’t it be nice if R had some helpful functions that would do all these tedious calculations at once? Something like `summary()` or `describe()`, perhaps? Why yes, yes it would. So much so that both of these functions exist. The `summary()` function is in the `base` package, so it comes with every installation of R. The `describe()` function is part of the `psych` package, which we loaded earlier in the chapter.

5.4.1 “Summarising” a variable

The `summary()` function is an easy thing to use, but a tricky thing to understand in full, since it’s a generic function (see Section 4.11). The basic idea behind the `summary()` function is that it prints out some useful information about whatever object (i.e., variable, as far as we’re concerned) you specify as the `object` argument. As a consequence, the behaviour of the `summary()` function differs quite dramatically depending on the class of the object that you give it. Let’s start by giving it a *numeric* object:

```
> summary( object = afl.margins )
   Min. 1st Qu. Median Mean 3rd Qu. Max.
0.00   12.75  30.50 35.30  50.50 116.00
```

For numeric variables, we get a whole bunch of useful descriptive statistics. It gives us the minimum and maximum values (i.e., the range), the first and third quartiles (25th and 75th percentiles; i.e., the IQR), the mean and the median. In other words, it gives us a pretty good collection of descriptive statistics related to the central tendency and the spread of the data.

Okay, what about if we feed it a logical vector instead? Let's say I want to know something about how many "blowouts" there were in the 2010 AFL season. I operationalise the concept of a blowout (see Chapter 2) as a game in which the winning margin exceeds 50 points. Let's create a logical variable `blowouts` in which the i -th element is `TRUE` if that game was a blowout according to my definition,

```
> blowouts <- afl.margins > 50
> blowouts
[1] TRUE FALSE TRUE FALSE FALSE FALSE FALSE TRUE FALSE FALSE
[11] FALSE TRUE FALSE FALSE TRUE FALSE FALSE FALSE FALSE TRUE
[21] FALSE FALSE FALSE FALSE FALSE FALSE TRUE FALSE TRUE
BLAH BLAH BLAH
```

So that's what the `blowouts` variable looks like. Now let's ask R for a `summary()`

```
> summary( object = blowouts )
  Mode   FALSE    TRUE    NA's
logical     132      44      0
```

In this context, the `summary()` function gives us a count of the number of `TRUE` values, the number of `FALSE` values, and the number of missing values (i.e., the `NAs`). Pretty reasonable behaviour.

Next, let's try to give it a factor. If you recall, I've defined the `afl.finalists` vector as a factor, so let's use that:

```
> summary( object = afl.finalists )
  Adelaide        Brisbane        Carlton        Collingwood
          26              25              26              28
  Essendon         Fitzroy        Fremantle        Geelong
          32              0                6              39
  Hawthorn        Melbourne    North Melbourne    Port Adelaide
          27              28              28              17
  Richmond        St Kilda        Sydney        West Coast
          6                24              26              38
  Western Bulldogs
          24
```

For factors, we get a frequency table, just like we got when we used the `table()` function. Interestingly, however, if we convert this to a character vector using the `as.character()` function (see Section 7.10), we don't get the same results:

```
> f2 <- as.character( afl.finalists )
> summary( object = f2 )
  Length   Class    Mode
  400    character  character
```

This is one of those situations I was referring to in Section 4.7, in which it is helpful to declare your nominal scale variable as a factor rather than a character vector. Because I've defined `afl.finalists` as a factor, R *knows* that it should treat it as a nominal scale variable, and so it gives you a much more detailed (and helpful) summary than it would have if I'd left it as a character vector.

5.4.2 “Summarising” a data frame

Okay what about data frames? When you pass a data frame to the `summary()` function, it produces a slightly condensed summary of each variable inside the data frame. To give you a sense of how this can

be useful, let's try this for a new data set, one that you've never seen before. The data is stored in the `clinicaltrial.Rdata` file, and we'll use it a lot in Chapter 14 (you can find a complete description of the data at the start of that chapter). Let's load it, and see what we've got:

```
> load( "clinicaltrial.Rdata" )
> who(TRUE)
-- Name --      -- Class --      -- Size --
clin.trial     data.frame     18 x 3
$drug          factor        18
$therapy       factor        18
$mood.gain    numeric       18
```

There's a single data frame called `clin.trial` which contains three variables, `drug`, `therapy` and `mood.gain`. Presumably then, this data is from a clinical trial of some kind, in which people were administered different drugs; and the researchers looked to see what the drugs did to their mood. Let's see if the `summary()` function sheds a little more light on this situation:

```
> summary( clin.trial )
   drug           therapy      mood.gain
placebo :6    no.therapy:9   Min.   :0.1000
anxitfree:6   CBT       :9   1st Qu.:0.4250
joyzepam:6    Median   :0.8500
                           Mean   :0.8833
                           3rd Qu.:1.3000
                           Max.   :1.8000
```

Evidently there were three drugs: a placebo, something called “anxitfree” and something called “joyzepam”; and there were 6 people administered each drug. There were 9 people treated using cognitive behavioural therapy (CBT) and 9 people who received no psychological treatment. And we can see from looking at the summary of the `mood.gain` variable that most people did show a mood gain (mean = .88), though without knowing what the scale is here it's hard to say much more than that. Still, that's not too bad. Overall, I feel that I learned something from that.

5.4.3 “Describing” a data frame

The `describe()` function (in the `psych` package) is a little different, and it's really only intended to be useful when your data are interval or ratio scale. Unlike the `summary()` function, it calculates the same descriptive statistics for any type of variable you give it. By default, these are:

- `var`. This is just an index: 1 for the first variable, 2 for the second variable, and so on.
- `n`. This is the sample size: more precisely, it's the number of non-missing values.
- `mean`. This is the sample mean (Section 5.1.1).
- `sd`. This is the (bias corrected) standard deviation (Section 5.2.5).
- `median`. The median (Section 5.1.3).
- `trimmed`. This is trimmed mean. By default it's the 10% trimmed mean (Section 5.1.6).
- `mad`. The median absolute deviation (Section 5.2.6).
- `min`. The minimum value.
- `max`. The maximum value.
- `range`. The range spanned by the data (Section 5.2.1).
- `skew`. The skewness (Section 5.3).
- `kurtosis`. The kurtosis (Section 5.3).
- `se`. The standard error of the mean (Chapter 10).

Notice that these descriptive statistics generally only make sense for data that are interval or ratio scale (usually encoded as numeric vectors). For nominal or ordinal variables (usually encoded as factors), most of these descriptive statistics are not all that useful. What the `describe()` function does is convert factors and logical variables to numeric vectors in order to do the calculations. These variables are marked with * and most of the time, the descriptive statistics for those variables won't make much sense. If you try to feed it a data frame that includes a character vector as a variable, it produces an error.

With those caveats in mind, let's use the `describe()` function to have a look at the `clin.trial` data frame. Here's what we get:

```
> describe( x = clin.trial )
      var n mean   sd median trimmed  mad min max range skew kurtosis    se
drug*     1 18 2.00 0.84    2.00   2.00 1.48 1.0 3.0    2.0 0.00    -1.66 0.20
therapy*  2 18 1.50 0.51    1.50   1.50 0.74 1.0 2.0    1.0 0.00    -2.11 0.12
mood.gain 3 18 0.88 0.53    0.85   0.88 0.67 0.1 1.8    1.7 0.13    -1.44 0.13
```

As you can see, the output for the asterisked variables is pretty meaningless, and should be ignored. However, for the `mood.gain` variable, there's a lot of useful information.

5.5

Descriptive statistics separately for each group

It is very commonly the case that you find yourself needing to look at descriptive statistics, broken down by some grouping variable. This is pretty easy to do in R, and there are three functions in particular that are worth knowing about: `by()`, `describeBy()` and `aggregate()`. Let's start with the `describeBy()` function, which is part of the `psych` package. The `describeBy()` function is very similar to the `describe()` function, except that it has an additional argument called `group` which specifies a grouping variable. For instance, let's say, I want to look at the descriptive statistics for the `clin.trial` data, broken down separately by `therapy` type. The command I would use here is:

```
> describeBy( x=clin.trial, group=clin.trial$therapy )

group: no.therapy
      var n mean   sd median trimmed  mad min max range skew kurtosis    se
drug*     1 9 2.00 0.87    2.00   2.00 1.48 1.0 3.0    2.0 0.00    -1.81 0.29
therapy*  2 9 1.00 0.00    1.00   1.00 0.00 1.0 1.0    0.0  NaN     NaN 0.00
mood.gain 3 9 0.72 0.59    0.72   0.72 0.44 0.1 1.7    1.6 0.51    -1.59 0.20
-----
group: CBT
      var n mean   sd median trimmed  mad min max range skew kurtosis    se
drug*     1 9 2.00 0.87    2.00   2.00 1.48 1.0 3.0    2.0 0.00    -1.81 0.29
therapy*  2 9 2.00 0.00    2.00   2.00 0.00 2.0 2.0    0.0  NaN     NaN 0.00
mood.gain 3 9 1.04 0.45    1.04   1.04 0.44 0.3 1.8    1.5 -0.03    -1.12 0.15
```

As you can see, the output is essentially identical to the output that the `describe()` function produce, except that the output now gives you means, standard deviations etc separately for the `CBT` group and the `no.therapy` group. Notice that, as before, the output displays asterisks for factor variables, in order to draw your attention to the fact that the descriptive statistics that it has calculated won't be very meaningful for those variables. Nevertheless, this command has given us some really useful descriptive statistics `mood.gain` variable, broken down as a function of `therapy`.

A somewhat more general solution is offered by the `by()` function. There are three arguments that you need to specify when using this function: the `data` argument specifies the data set, the `INDICES`

argument specifies the grouping variable, and the `FUN` argument specifies the name of a function that you want to apply separately to each group. To give a sense of how powerful this is, you can reproduce the `describeBy()` function by using a command like this:

```
> by( data=clin.trial, INDICES=clin.trial$therapy, FUN=describe )
```

This will produce the exact same output as the command shown earlier. However, there's nothing special about the `describe()` function. You could just as easily use the `by()` function in conjunction with the `summary()` function. For example:

```
> by( data=clin.trial, INDICES=clin.trial$therapy, FUN=summary )
```

```
clin.trial$therapy: no.therapy
  drug      therapy   mood.gain
placebo :3  no.therapy:9   Min.   :0.1000
anxitfree:3    CBT       :0   1st Qu.:0.3000
joyzepam:3          Median  :0.5000
                           Mean   :0.7222
                           3rd Qu.:1.3000
                           Max.   :1.7000
-----
clin.trial$therapy: CBT
  drug      therapy   mood.gain
placebo :3  no.therapy:0   Min.   :0.300
anxitfree:3    CBT       :9   1st Qu.:0.800
joyzepam:3          Median  :1.100
                           Mean   :1.044
                           3rd Qu.:1.300
                           Max.   :1.800
```

Again, this output is pretty easy to interpret. It's the output of the `summary()` function, applied separately to `CBT` group and the `no.therapy` group. For the two factors (`drug` and `therapy`) it prints out a frequency table, whereas for the numeric variable (`mood.gain`) it prints out the range, interquartile range, mean and median.

What if you have multiple grouping variables? Suppose, for example, you would like to look at the average mood gain separately for all possible combinations of drug and therapy. It is actually possible to do this using the `by()` and `describeBy()` functions, but I usually find it more convenient to use the `aggregate()` function in this situation. There are again three arguments that you need to specify. The `formula` argument is used to indicate which variable you want to analyse, and which variables are used to specify the groups. For instance, if you want to look at `mood.gain` separately for each possible combination of `drug` and `therapy`, the formula you want is `mood.gain ~ drug + therapy`. The `data` argument is used to specify the data frame containing all the data, and the `FUN` argument is used to indicate what function you want to calculate for each group (e.g., the `mean`). So, to obtain group means, use this command:

```
> aggregate( formula = mood.gain ~ drug + therapy, # mood.gain by drug/therapy combination
+               data = clin.trial,                      # data is in the clin.trial data frame
+               FUN = mean                                # print out group means
+ )
  drug      therapy mood.gain
1 placebo no.therapy 0.300000
2 anxitfree no.therapy 0.400000
3 joyzepam no.therapy 1.466667
4 placebo      CBT 0.600000
5 anxitfree      CBT 1.033333
6 joyzepam      CBT 1.500000
```

or, alternatively, if you want to calculate the standard deviations for each group, you would use the following command (argument names omitted this time):

```
> aggregate( mood.gain ~ drug + therapy, clin.trial, sd )
   drug      therapy mood.gain
1 placebo no.therapy 0.2000000
2 anxifree no.therapy 0.2000000
3 joyzepam no.therapy 0.2081666
4 placebo      CBT 0.3000000
5 anxifree      CBT 0.2081666
6 joyzepam      CBT 0.2645751
```

5.6 _____

Standard scores

Suppose my friend is putting together a new questionnaire intended to measure “grumpiness”. The survey has 50 questions, which you can answer in a grumpy way or not. Across a big sample (hypothetically, let’s imagine a million people or so!) the data are fairly normally distributed, with the mean grumpiness score being 17 out of 50 questions answered in a grumpy way, and the standard deviation is 5. In contrast, when I take the questionnaire, I answer 35 out of 50 questions in a grumpy way. So, how grumpy am I? One way to think about would be to say that I have grumpiness of 35/50, so you might say that I’m 70% grumpy. But that’s a bit weird, when you think about it. If my friend had phrased her questions a bit differently, people might have answered them in a different way, so the overall distribution of answers could easily move up or down depending on the precise way in which the questions were asked. So, I’m only 70% grumpy *with respect to this set of survey questions*. Even if it’s a very good questionnaire, this isn’t very a informative statement.

A simpler way around this is to describe my grumpiness by comparing me to other people. Shockingly, out of my friend’s sample of 1,000,000 people, only 159 people were as grumpy as me (that’s not at all unrealistic, frankly), suggesting that I’m in the top 0.016% of people for grumpiness. This makes much more sense than trying to interpret the raw data. This idea – that we should describe my grumpiness in terms of the overall distribution of the grumpiness of humans – is the qualitative idea that standardisation attempts to get at. One way to do this is to do exactly what I just did, and describe everything in terms of percentiles. However, the problem with doing this is that “it’s lonely at the top”. Suppose that my friend had only collected a sample of 1000 people (still a pretty big sample for the purposes of testing a new questionnaire, I’d like to add), and this time gotten a mean of 16 out of 50 with a standard deviation of 5, let’s say. The problem is that almost certainly, not a single person in that sample would be as grumpy as me.

However, all is not lost. A different approach is to convert my grumpiness score into a **standard score**, also referred to as a *z-score*. The standard score is defined as the number of standard deviations above the mean that my grumpiness score lies. To phrase it in “pseudo-maths” the standard score is calculated like this:

$$\text{standard score} = \frac{\text{raw score} - \text{mean}}{\text{standard deviation}}$$

In actual maths, the equation for the *z-score* is

$$z_i = \frac{X_i - \bar{X}}{\hat{\sigma}}$$

So, going back to the grumpiness data, we can now transform Dan’s raw grumpiness into a standardised

grumpiness score.¹³ If the mean is 17 and the standard deviation is 5 then my standardised grumpiness score would be¹⁴

$$z = \frac{35 - 17}{5} = 3.6$$

To interpret this value, recall the rough heuristic that I provided in Section 5.2.5, in which I noted that 99.7% of values are expected to lie within 3 standard deviations of the mean. So the fact that my grumpiness corresponds to a z score of 3.6 indicates that I'm very grumpy indeed. Later on, in Section 9.4, I'll introduce a function called `pnorm()` that allows us to be a bit more precise than this. Specifically, it allows us to calculate a theoretical percentile rank for my grumpiness, as follows:

```
> pnorm( 3.6 )
[1] 0.9998409
```

At this stage, this command doesn't make too much sense, but don't worry too much about it. It's not important for now. But the output is fairly straightforward: it suggests that I'm grumpier than 99.98% of people. Sounds about right.

In addition to allowing you to interpret a raw score in relation to a larger population (and thereby allowing you to make sense of variables that lie on arbitrary scales), standard scores serve a second useful function. Standard scores can be compared to one another in situations where the raw scores can't. Suppose, for instance, my friend also had another questionnaire that measured extraversion using a 24 items questionnaire. The overall mean for this measure turns out to be 13 with standard deviation 4; and I scored a 2. As you can imagine, it doesn't make a lot of sense to try to compare my raw score of 2 on the extraversion questionnaire to my raw score of 35 on the grumpiness questionnaire. The raw scores for the two variables are "about" fundamentally different things, so this would be like comparing apples to oranges.

What about the standard scores? Well, this is a little different. If we calculate the standard scores, we get $z = (35 - 17)/5 = 3.6$ for grumpiness and $z = (2 - 13)/4 = -2.75$ for extraversion. These two numbers *can* be compared to each other.¹⁵ I'm much less extraverted than most people ($z = -2.75$) and much grumpier than most people ($z = 3.6$): but the extent of my unusualness is much more extreme for grumpiness (since 3.6 is a bigger number than 2.75). Because each standardised score is a statement about where an observation falls *relative to its own population*, it is possible to compare standardised scores across completely different variables.

5.7

Correlations

Up to this point we have focused entirely on how to construct descriptive statistics for a single variable. What we haven't done is talked about how to describe the relationships *between* variables in the data. To do that, we want to talk mostly about the **correlation** between variables. But first, we need some data.

¹³I haven't discussed how to compute z -scores, explicitly, but you can probably guess. For a variable `x`, the simplest way is to use a command like `(x - mean(x)) / sd(x)`. There's also a fancier function called `scale()` that you can use, but it relies on somewhat more complicated R concepts that I haven't explained yet.

¹⁴Technically, because I'm calculating means and standard deviations from a sample of data, but want to talk about my grumpiness relative to a population, what I'm actually doing is *estimating* a z score. However, since we haven't talked about estimation yet (see Chapter 10) I think it's best to ignore this subtlety, especially as it makes very little difference to our calculations.

¹⁵Though some caution is usually warranted. It's not always the case that one standard deviation on variable A corresponds to the same "kind" of thing as one standard deviation on variable B. Use common sense when trying to determine whether or not the z scores of two variables can be meaningfully compared.

Table 5.1: Descriptive statistics for the parenthood data.

variable	min	max	mean	median	std. dev	IQR
Dan's grumpiness	41	91	63.71	62	10.05	14
Dan's hours slept	4.84	9.00	6.97	7.03	1.02	1.45
Dan's son's hours slept	3.25	12.07	8.05	7.95	2.07	3.21

5.7.1 The data

After spending so much time looking at the AFL data, I'm starting to get bored with sports. Instead, let's turn to a topic close to every parent's heart: sleep. The following data set is fictitious, but based on real events. Suppose I'm curious to find out how much my infant son's sleeping habits affect my mood. Let's say that I can rate my grumpiness very precisely, on a scale from 0 (not at all grumpy) to 100 (grumpy as a very, very grumpy old man). And, lets also assume that I've been measuring my grumpiness, my sleeping patterns and my son's sleeping patterns for quite some time now. Let's say, for 100 days. And, being a nerd, I've saved the data as a file called `parenthood.Rdata`. If we load the data...

```
> load("parenthood.Rdata")
> who(TRUE)
-- Name --    -- Class --    -- Size --
parenthood   data.frame   100 x 4
$dan.sleep   numeric     100
$baby.sleep  numeric     100
$dan.grump   numeric     100
$day         integer     100
```

... we see that the file contains a single data frame called `parenthood`, which contains four variables `dan.sleep`, `baby.sleep`, `dan.grump` and `day`. If we print out the data, here's what we get:

```
> parenthood
  dan.sleep baby.sleep dan.grump day
1      7.59     10.18       56    1
2      7.91     11.66       60    2
3      5.14      7.92       82    3
4      7.71      9.61       55    4

BLAH BLAH BLAH

98      5.38      6.97       82   98
99      7.02      6.56       55   99
100     6.45      7.93       74  100
```

Next, I'll calculate some basic descriptive statistics:

```
> describe(parenthood)
      var   n   mean     sd median trimmed   mad   min   max
dan.sleep  1 100  6.97  1.02    7.03   7.00  1.09  4.84  9.00
baby.sleep 2 100  8.05  2.07    7.95   8.05  2.33  3.25 12.07
dan.grump  3 100 63.71 10.05   62.00  63.16  9.64 41.00 91.00
day        4 100 50.50 29.01   50.50  50.50 37.06  1.00 100.00
```

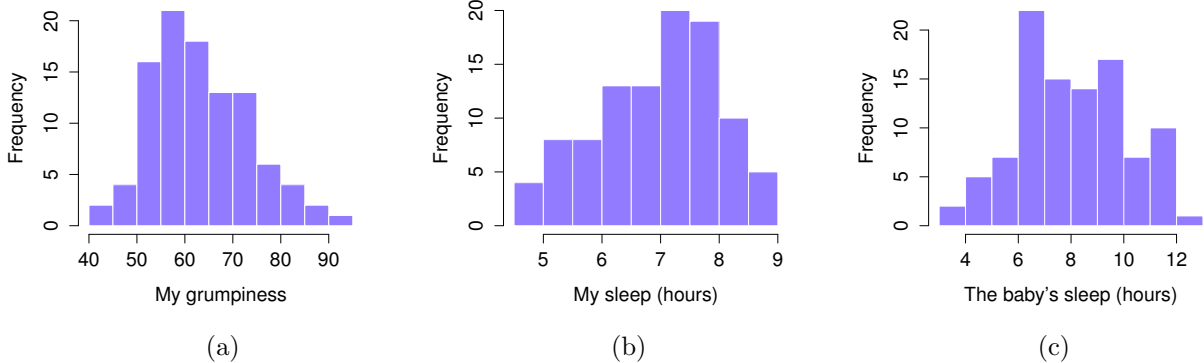


Figure 5.6: Histograms for the three interesting variables in the `parenthood` data set.

	range	skew	kurtosis	se
dan.sleep	4.16	-0.29	-0.65	0.10
baby.sleep	8.82	-0.02	-0.61	0.21
dan.grump	50.00	0.43	-0.04	1.00
day	99.00	0.00	-1.20	2.90

Finally, to give a graphical depiction of what each of the three interesting variables looks like, Figure 5.6 plots histograms. One thing to note: just because R can calculate dozens of different statistics doesn't mean you should report all of them. If I were writing this up for a report, I'd probably pick out those statistics that are of most interest to me (and to my readership), and then put them into a nice, simple table like the one in Table 5.1.¹⁶ Notice that when I put it into a table, I gave everything "human readable" names. This is always good practice. Notice also that I'm not getting enough sleep. This isn't good practice, but other parents tell me that it's standard practice.

5.7.2 The strength and direction of a relationship

We can draw scatterplots to give us a general sense of how closely related two variables are. Ideally though, we might want to say a bit more about it than that. For instance, let's compare the relationship between `dan.sleep` and `dan.grump` (Figure 5.7, left) with that between `baby.sleep` and `dan.grump` (Figure 5.7, right). When looking at these two plots side by side, it's clear that the relationship is *qualitatively* the same in both cases: more sleep equals less grump! However, it's also pretty obvious that the relationship between `dan.sleep` and `dan.grump` is *stronger* than the relationship between `baby.sleep` and `dan.grump`. The plot on the left is "neater" than the one on the right. What it feels like is that if you want to predict what my mood is, it'd help you a little bit to know how many hours my son slept, but it'd be more helpful to know how many hours I slept.

In contrast, let's consider the two scatterplots shown in Figure 5.8. If we compare the scatterplot of "`baby.sleep v dan.grump`" (left) to the scatterplot of "`baby.sleep v dan.sleep`" (right), the overall strength of the relationship is the same, but the direction is different. That is, if my son sleeps more,

¹⁶Actually, even that table is more than I'd bother with. In practice most people pick *one* measure of central tendency, and *one* measure of variability only.

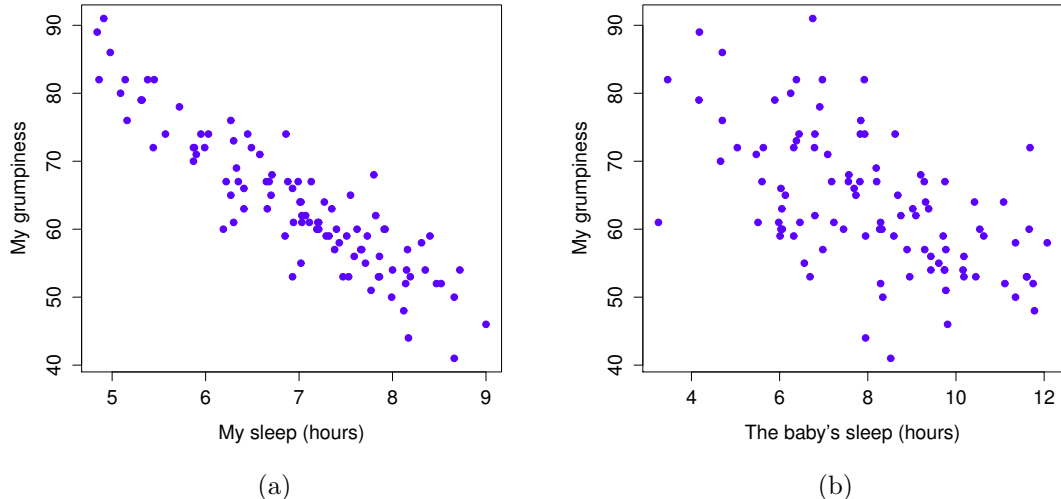


Figure 5.7: Scatterplots showing the relationship between `dan.sleep` and `dan.grump` (left) and the relationship between `baby.sleep` and `dan.grump` (right).

I get *more* sleep (positive relationship, right hand side), but if he sleeps more then I get *less* grumpy (negative relationship, left hand side).

5.7.3 The correlation coefficient

We can make these ideas a bit more explicit by introducing the idea of a **correlation coefficient** (or, more specifically, Pearson’s correlation coefficient), which is traditionally denoted by r . The correlation coefficient between two variables X and Y (sometimes denoted r_{XY}), which we’ll define more precisely in the next section, is a measure that varies from -1 to 1 . When $r = -1$ it means that we have a perfect negative relationship, and when $r = 1$ it means we have a perfect positive relationship. When $r = 0$, there’s no relationship at all. If you look at Figure 5.9, you can see several plots showing what different correlations look like.

The formula for the Pearson’s correlation coefficient can be written in several different ways. I think the simplest way to write down the formula is to break it into two steps. Firstly, let’s introduce the idea of a **covariance**. The covariance between two variables X and Y is a generalisation of the notion of the variance; it’s a mathematically simple way of describing the relationship between two variables that isn’t terribly informative to humans:

$$\text{Cov}(X, Y) = \frac{1}{N-1} \sum_{i=1}^N (X_i - \bar{X})(Y_i - \bar{Y})$$

Because we’re multiplying (i.e., taking the “product” of) a quantity that depends on X by a quantity that depends on Y and then averaging¹⁷, you can think of the formula for the covariance as an “average cross product” between X and Y . The covariance has the nice property that, if X and Y are entirely unrelated, then the covariance is exactly zero. If the relationship between them is positive (in the

¹⁷Just like we saw with the variance and the standard deviation, in practice we divide by $N - 1$ rather than N .

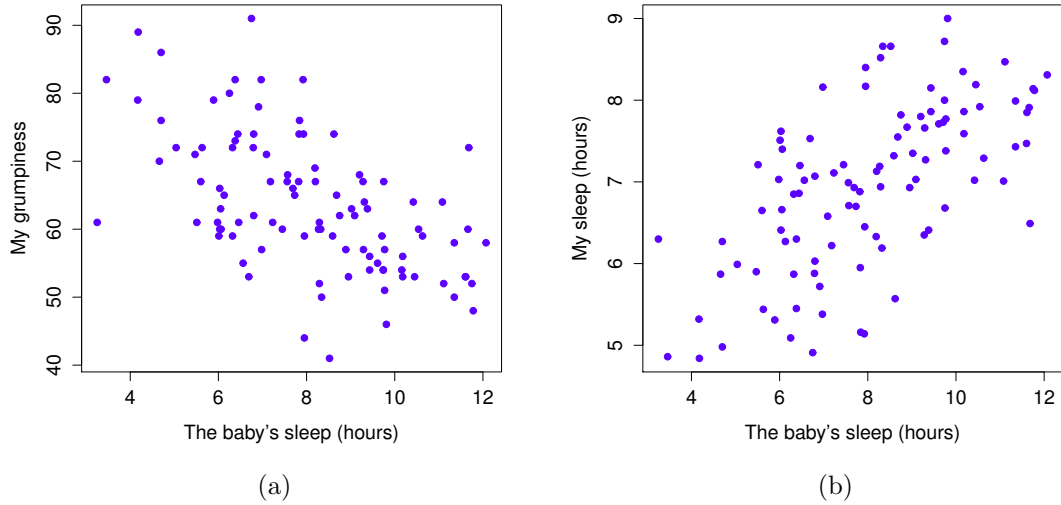


Figure 5.8: Scatterplots showing the relationship between `baby.sleep` and `dan.grump` (left), as compared to the relationship between `baby.sleep` and `dan.sleep` (right).

sense shown in Figure 5.9) then the covariance is also positive; and if the relationship is negative then the covariance is also negative. In other words, the covariance captures the basic qualitative idea of correlation. Unfortunately, the raw magnitude of the covariance isn't easy to interpret: it depends on the units in which X and Y are expressed, and worse yet, the actual units that the covariance itself is expressed in are really weird. For instance, if X refers to the `dan.sleep` variable (units: hours) and Y refers to the `dan.grump` variable (units: grumps), then the units for their covariance are “hours \times grumps”. And I have no freaking idea what that would even mean.

The Pearson correlation coefficient r fixes this interpretation problem by standardising the covariance, in pretty much the exact same way that the z -score standardises a raw score: by dividing by the standard deviation. However, because we have two variables that contribute to the covariance, the standardisation only works if we divide by both standard deviations.¹⁸ In other words, the correlation between X and Y can be written as follows:

$$r_{XY} = \frac{\text{Cov}(X, Y)}{\hat{\sigma}_X \hat{\sigma}_Y}$$

By doing this standardisation, not only do we keep all of the nice properties of the covariance discussed earlier, but the actual values of r are on a meaningful scale: $r = 1$ implies a perfect positive relationship, and $r = -1$ implies a perfect negative relationship. I'll expand a little more on this point later, in Section 5.7.5. But before I do, let's look at how to calculate correlations in R.

5.7.4 Calculating correlations in R

Calculating correlations in R can be done using the `cor()` command. The simplest way to use the command is to specify two input arguments `x` and `y`, each one corresponding to one of the variables. The

¹⁸This is an oversimplification, but it'll do for our purposes.

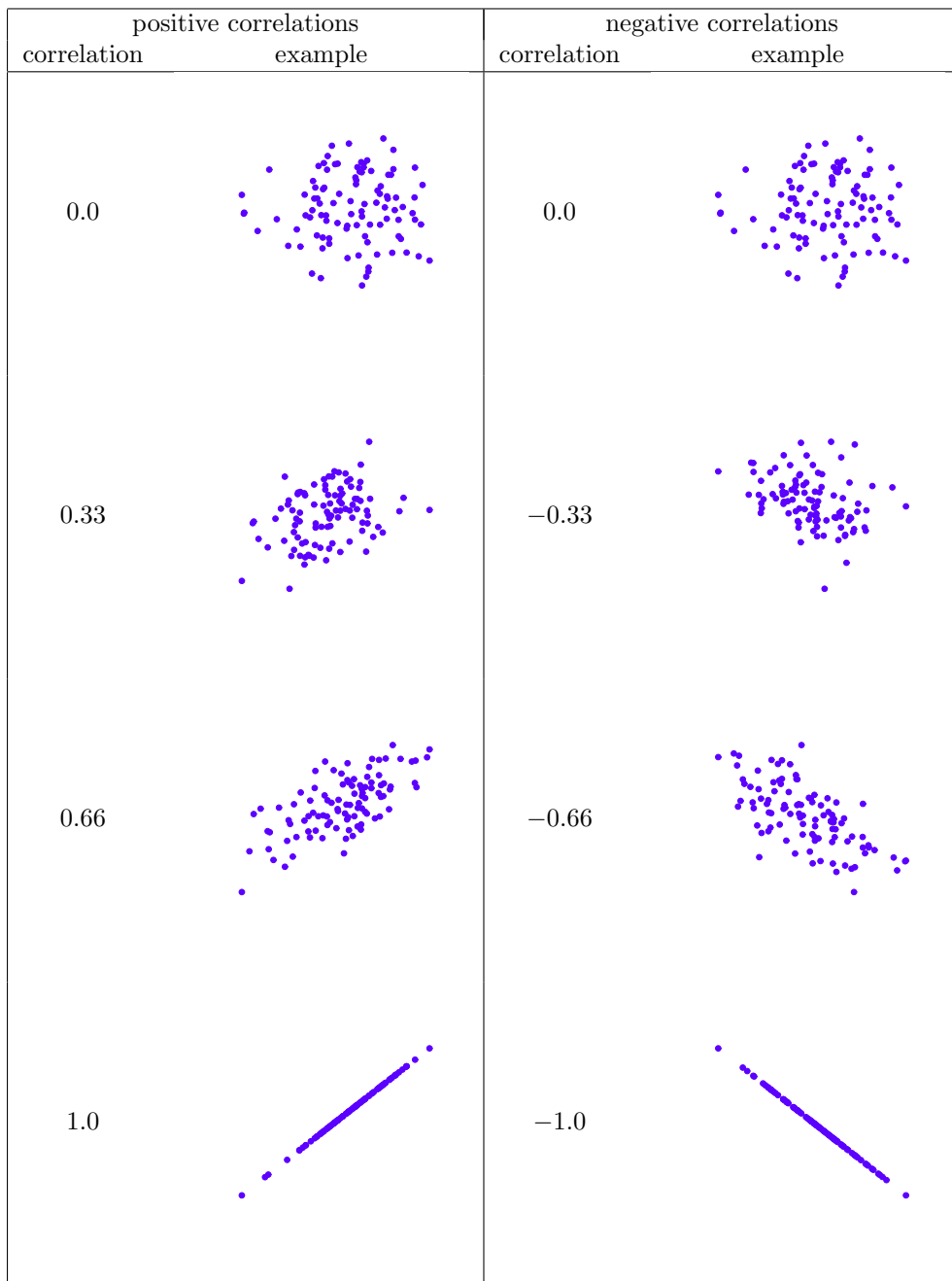


Figure 5.9: Illustration of the effect of varying the strength and direction of a correlation. In the left hand column, the correlations are 0, .33, .66 and 1. In the right hand column, the correlations are 0, -.33, -.66 and -1.

following extract illustrates the basic usage of the function:¹⁹

```
> cor( x = parenthood$dan.sleep, y = parenthood$dan.grump )
[1] -0.903384
```

However, the `cor()` function is a bit more powerful than this simple example suggests. For example, you can also calculate a complete “correlation matrix”, between all pairs of variables in the data frame.²⁰

```
> # correlate all pairs of variables in "parenthood":
> cor( x = parenthood )
      dan.sleep  baby.sleep  dan.grump       day
dan.sleep   1.00000000  0.62794934 -0.90338404 -0.09840768
baby.sleep  0.62794934  1.00000000 -0.56596373 -0.01043394
dan.grump  -0.90338404 -0.56596373  1.00000000  0.07647926
day        -0.09840768 -0.01043394  0.07647926  1.00000000
```

5.7.5 Interpreting a correlation

Naturally, in real life you don’t see many correlations of 1. So how should you interpret a correlation of, say $r = .4$? The honest answer is that it really depends on what you want to use the data for, and on how strong the correlations in your field tend to be. A friend of mine in engineering once argued that any correlation less than .95 is completely useless (I think he was exaggerating, even for engineering). On the other hand there are real cases – even in psychology – where you should really expect correlations that strong. For instance, one of the benchmark data sets used to test theories of how people judge similarities is so clean that any theory that can’t achieve a correlation of at least .9 really isn’t deemed to be successful. However, when looking for (say) elementary correlates of intelligence (e.g., inspection time, response time), if you get a correlation above .3 you’re doing very very well. In short, the interpretation of a correlation depends a lot on the context. That said, the rough guide in Table 5.2 is pretty typical.

However, something that can never be stressed enough is that you should *always* look at the scatterplot before attaching any interpretation to the data. A correlation might not mean what you think it means. The classic illustration of this is “Anscombe’s Quartet” (Anscombe, 1973), which is a collection of four data sets. Each data set has two variables, an X and a Y . For all four data sets the mean value for X is 9 and the mean for Y is 7.5. The standard deviations for all X variables are almost identical, as are those for the Y variables. And in each case the correlation between X and Y is $r = 0.816$. You can verify this yourself, since I happen to have saved the in a file²¹ called `anscombesquartet.Rdata`. The commands would be:

```
> load( "anscombesquartet.Rdata" )
> cor( X1, Y1 )
[1] 0.8164205
> cor( X2, Y2 )
[1] 0.8162365
```

and so on. You’d think that these four data sets would look pretty similar to one another. They do not. If we draw scatterplots of X against Y for all four variables, as shown in Figure 5.10 we see that all four

¹⁹If you are reading this after having already completed Chapter 11 you might be wondering about hypothesis tests for correlations. R has a function called `cor.test()` that runs a hypothesis test for a single correlation, and the `psych` package contains a version called `corr.test()` that can run tests for every correlation in a correlation matrix; hypothesis tests for correlations are discussed in more detail in Section 15.5.4.

²⁰An alternative usage of `cor()` is to correlate one set of variables with another subset of variables. If `x` and `y` are both data frames with the same number of rows, then `cor(x = x, y = Y)` will produce a correlation matrix that correlates all variables in `x` with all variables in `y`.

²¹It’s also stored as a dataset that comes distributed with R. Just type `anscombe` at the command prompt.

Table 5.2: A rough guide to interpreting correlations. Note that I say a *rough* guide. There aren't hard and fast rules for what counts as strong or weak relationships. It depends on the context.

Correlation	Strength	Direction
-1.0 to -0.9	Very strong	Negative
-0.9 to -0.7	Strong	Negative
-0.7 to -0.4	Moderate	Negative
-0.4 to -0.2	Weak	Negative
-0.2 to 0	Negligible	Negative
0 to 0.2	Negligible	Positive
0.2 to 0.4	Weak	Positive
0.4 to 0.7	Moderate	Positive
0.7 to 0.9	Strong	Positive
0.9 to 1.0	Very strong	Positive

of these are *spectacularly* different to each other. The lesson here, which so very many people seem to forget in real life is “*always graph your raw data*”. This will be the focus of Chapter 6.

5.7.6 Spearman's rank correlations

The Pearson correlation coefficient is useful for a lot of things, but it does have shortcomings. One issue in particular stands out: what it actually measures is the strength of the *linear* relationship between two variables. In other words, what it gives you is a measure of the extent to which the data all tend to fall on a single, perfectly straight line. Often, this is a pretty good approximation to what we mean when we say “relationship”, and so the Pearson correlation is a good thing to calculate. Sometimes, it isn’t.

One very common situation where the Pearson correlation isn’t quite the right thing to use arises when an increase in one variable X really is reflected in an increase in another variable Y , but the nature of the relationship isn’t necessarily linear. An example of this might be the relationship between effort and reward when studying for an exam. If you put in zero effort (X) into learning a subject, then you should expect a grade of 0% (Y). However, a little bit of effort will cause a *massive* improvement: just turning up to lectures means that you learn a fair bit, and if you just turn up to classes, and scribble a few things down so your grade might rise to 35%, all without a lot of effort. However, you just don’t get the same effect at the other end of the scale. As everyone knows, it takes a *lot* more effort to get a grade of 90% than it takes to get a grade of 55%. What this means is that, if I’ve got data looking at study effort and grades, there’s a pretty good chance that Pearson correlations will be misleading.

To illustrate, consider the data plotted in Figure 5.11, showing the relationship between hours worked and grade received for 10 students taking some class. The curious thing about this – highly fictitious – data set is that increasing your effort *always* increases your grade. It might be by a lot or it might be by a little, but increasing effort will never decrease your grade. The data are stored in `effort.Rdata`:

```
> load("effort.Rdata")
> who(TRUE)
-- Name --  -- Class --  -- Size --
effort      data.frame    10 x 2
$hours      numeric       10
$grade      numeric       10
```

The raw data look like this:

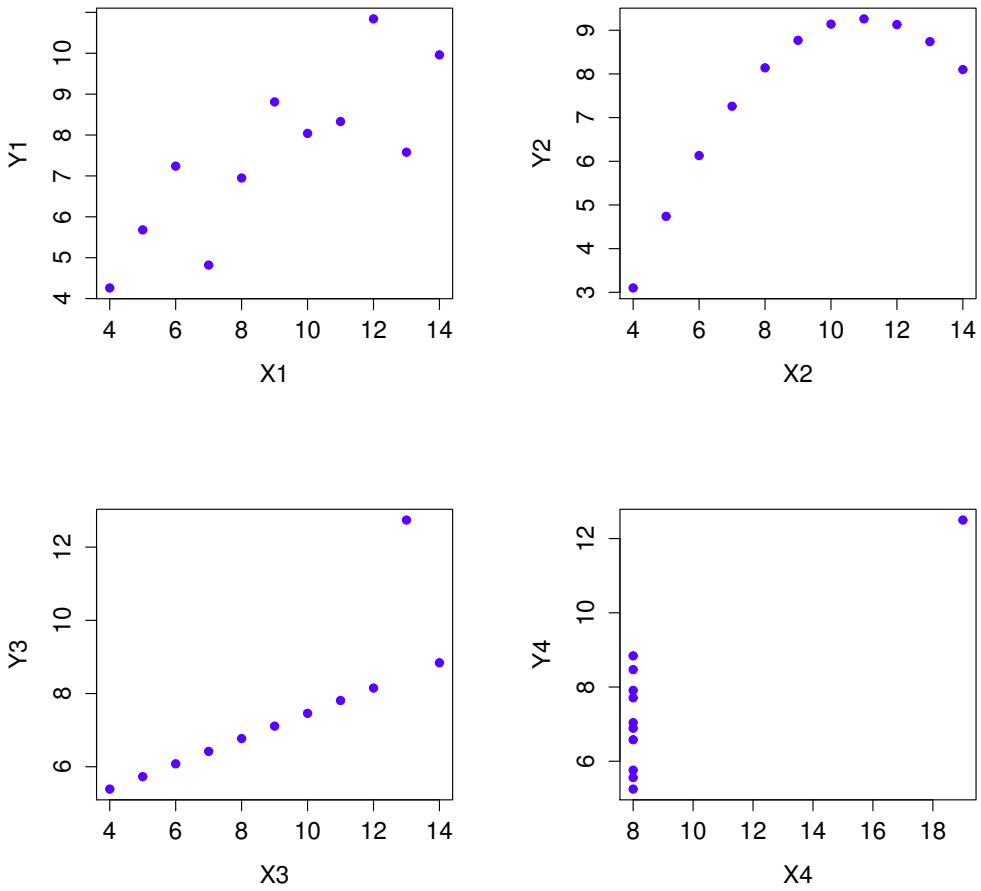


Figure 5.10: Anscombe's quartet. All four of these data sets have a Pearson correlation of $r = .816$, but they are qualitatively different from one another.

```
> effort
  hours grade
1      2    13
2     76    91
3     40    79
4      6    14
5     16    21
6     28    74
7     27    47
8     59    85
9     46    84
10    68    88
```

If we run a standard Pearson correlation, it shows a strong relationship between hours worked and grade received,

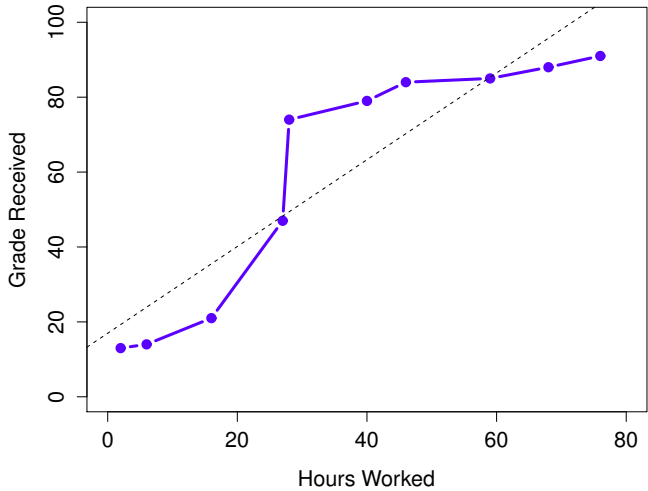


Figure 5.11: The relationship between hours worked and grade received, for a toy data set consisting of only 10 students (each circle corresponds to one student). The dashed line through the middle shows the linear relationship between the two variables. This produces a strong Pearson correlation of $r = .91$. However, the interesting thing to note here is that there's actually a perfect monotonic relationship between the two variables: in this toy example at least, increasing the hours worked always increases the grade received, as illustrated by the solid line. This is reflected in a Spearman correlation of $\rho = 1$. With such a small data set, however, it's an open question as to which version better describes the actual relationship involved.

```
> cor( effort$hours, effort$grade )
[1] 0.909402
```

but this doesn't actually capture the observation that increasing hours worked *always* increases the grade. There's a sense here in which we want to be able to say that the correlation is *perfect* but for a somewhat different notion of what a "relationship" is. What we're looking for is something that captures the fact that there is a perfect **ordinal relationship** here. That is, if student 1 works more hours than student 2, then we can guarantee that student 1 will get the better grade. That's not what a correlation of $r = .91$ says at all.

How should we address this? Actually, it's really easy: if we're looking for ordinal relationships, all we have to do is treat the data as if it were ordinal scale! So, instead of measuring effort in terms of "hours worked", let's rank all 10 of our students in order of hours worked. That is, student 1 did the least work out of anyone (2 hours) so they get the lowest rank (rank = 1). Student 4 was the next laziest, putting in only 6 hours of work in over the whole semester, so they get the next lowest rank (rank = 2). Notice that I'm using "rank = 1" to mean "low rank". Sometimes in everyday language we talk about "rank = 1" to mean "top rank" rather than "bottom rank". So be careful: you can rank "from smallest value to largest value" (i.e., small equals rank 1) or you can rank "from largest value to smallest value" (i.e., large equals rank 1). In this case, I'm ranking from smallest to largest, because that's the default way that R does it. But in real life, it's really easy to forget which way you set things up, so you have to put a bit of effort into remembering!

Okay, so let's have a look at our students when we rank them from worst to best in terms of effort and reward:

	rank (hours worked)	rank (grade received)
student 1	1	1
student 2	10	10
student 3	6	6
student 4	2	2
student 5	3	3
student 6	5	5
student 7	4	4
student 8	8	8
student 9	7	7
student 10	9	9

Hm. These are *identical*. The student who put in the most effort got the best grade, the student with the least effort got the worst grade, etc. We can get R to construct these rankings using the `rank()` function, like this:

```
> hours.rank <- rank( effort$hours )    # rank students by hours worked
> grade.rank <- rank( effort$grade )    # rank students by grade received
```

As the table above shows, these two rankings are identical, so if we now correlate them we get a perfect relationship:

```
> cor( hours.rank, grade.rank )
[1] 1
```

What we've just re-invented is **Spearman's rank order correlation**, usually denoted ρ to distinguish it from the Pearson correlation r . We can calculate Spearman's ρ using R in two different ways. Firstly we could do it the way I just showed, using the `rank()` function to construct the rankings, and then calculate the Pearson correlation on these ranks. However, that's way too much effort to do every time. It's much easier to just specify the `method` argument of the `cor()` function.

```
> cor( effort$hours, effort$grade, method = "spearman")
[1] 1
```

The default value of the `method` argument is "`pearson`", which is why we didn't have to specify it earlier on when we were doing Pearson correlations.

5.8

Handling missing values

There's one last topic that I want to discuss briefly in this chapter, and that's the issue of **missing data**. Real data sets very frequently turn out to have missing values: perhaps someone forgot to fill in a particular survey question, for instance. Missing data can be the source of a lot of tricky issues, most of which I'm going to gloss over. However, at a minimum, you need to understand the basics of handling missing data in R.

5.8.1 The single variable case

Let's start with the simplest case, in which you're trying to calculate descriptive statistics for a single variable which has missing data. In R, this means that there will be `NA` values in your data vector. Let's create a variable like that:

```
> partial <- c(10, 20, NA, 30)
```

Let's assume that you want to calculate the mean of this variable. By default, R assumes that you want to calculate the mean using all four elements of this vector, which is probably the safest thing for a dumb automaton to do, but it's rarely what you actually want. Why not? Well, remember that the basic interpretation of `NA` is "I don't know what this number is". This means that `1 + NA = NA`: if I add 1 to some number that I don't know (i.e., the `NA`) then the answer is *also* a number that I don't know. As a consequence, if you don't explicitly tell R to ignore the `NA` values, and the data set does have missing values, then the output will itself be a missing value. If I try to calculate the mean of the `partial` vector, without doing anything about the missing value, here's what happens:

```
> mean( x = partial )
[1] NA
```

Technically correct, but deeply unhelpful.

To fix this, all of the descriptive statistics functions that I've discussed in this chapter (with the exception of `cor()` which is a special case I'll discuss below) have an optional argument called `na.rm`, which is shorthand for "remove NA values". By default, `na.rm = FALSE`, so R does nothing about the missing data problem. Let's try setting `na.rm = TRUE` and see what happens:

When calculating sums and means when missing data are present (i.e., when there are `NA` values) there's actually an additional argument to the function that you should be aware of. This argument is called `na.rm`, and is a logical value indicating whether R should ignore (or "remove") the missing data for the purposes of doing the calculations. By default, R assumes that you want to keep the missing values, so unless you say otherwise it will set `na.rm = FALSE`. However, R assumes that `1 + NA = NA`: if I add 1 to some number that I don't know (i.e., the `NA`) then the answer is *also* a number that I don't know. As a consequence, if you don't explicitly tell R to ignore the `NA` values, and the data set does have missing values, then the output will itself be a missing value. This is illustrated in the following extract:

```
> mean( x = partial, na.rm = TRUE )
[1] 20
```

Notice that the mean is `20` (i.e., `60 / 3`) and *not* `15`. When R ignores a `NA` value, it genuinely ignores it. In effect, the calculation above is identical to what you'd get if you asked for the mean of the three-element vector `c(10, 20, 30)`.

As indicated above, this isn't unique to the `mean()` function. Pretty much all of the other functions that I've talked about in this chapter have an `na.rm` argument that indicates whether it should ignore missing values. However, its behaviour is the same for all these functions, so I won't waste everyone's time by demonstrating it separately for each one.

5.8.2 Missing values in pairwise calculations

I mentioned earlier that the `cor()` function is a special case. It doesn't have an `na.rm` argument, because the story becomes a lot more complicated when more than one variable is involved. What it

does have is an argument called `use` which does roughly the same thing, but you need to think little more carefully about what you want this time. To illustrate the issues, let's open up a data set that has missing values, `parenthood2.Rdata`. This file contains the same data as the original parenthood data, but with some values deleted. It contains a single data frame, `parenthood2`:

```
> load("parenthood2.Rdata")
> print(parenthood2)
  dan.sleep baby.sleep dan.grump day
1      7.59        NA      56   1
2      7.91     11.66      60   2
3      5.14      7.92      82   3
4      7.71      9.61      55   4
5      6.68      9.75      NA   5
6      5.99      5.04      72   6
BLAH BLAH BLAH
```

If I calculate my descriptive statistics using the `describe()` function

```
> describe(parenthood2)
      var   n   mean     sd median trimmed   mad   min   max   BLAH
dan.sleep    1  91  6.98  1.02    7.03    7.02  1.13  4.84  9.00   BLAH
baby.sleep   2  89  8.11  2.05    8.20    8.13  2.28  3.25 12.07   BLAH
dan.grump    3  92 63.15  9.85   61.00   62.66 10.38 41.00 89.00   BLAH
day         4 100 50.50 29.01   50.50   50.50 37.06  1.00 100.00   BLAH
```

we can see from the `n` column that there are 9 missing values for `dan.sleep`, 11 missing values for `baby.sleep` and 8 missing values for `dan.grump`.²² Suppose what I would like is a correlation matrix. And let's also suppose that I don't bother to tell R how to handle those missing values. Here's what happens:

```
> cor(parenthood2)
      dan.sleep baby.sleep dan.grump day
dan.sleep       1        NA       NA   NA
baby.sleep      NA       1        NA   NA
dan.grump      NA       NA       1   NA
day            NA       NA       NA   1
```

Annoying, but it kind of makes sense. If I don't *know* what some of the values of `dan.sleep` and `baby.sleep` actually are, then I can't possibly *know* what the correlation between these two variables is either, since the formula for the correlation coefficient makes use of every single observation in the data set. Once again, it makes sense: it's just not particularly *helpful*.

To make R behave more sensibly in this situation, you need to specify the `use` argument to the `cor()` function. There are several different values that you can specify for this, but the two that we care most about in practice tend to be `"complete.obs"` and `"pairwise.complete.obs"`. If we specify `use = "complete.obs"`, R will completely ignore all cases (i.e., all rows in our `parenthood2` data frame) that have any missing values at all. So, for instance, if you look back at the extract earlier when I used the `head()` function, notice that observation 1 (i.e., day 1) of the `parenthood2` data set is missing the value for `baby.sleep`, but is otherwise complete? Well, if you choose `use = "complete.obs"` R will ignore that row completely: that is, even when it's trying to calculate the correlation between `dan.sleep` and `dan.grump`, observation 1 will be ignored, because the value of `baby.sleep` is missing for that observation. Here's what we get:

²²It's worth noting that, even though we have missing data for each of these variables, the output doesn't contain any `NA` values. This is because, while `describe()` also has an `na.rm` argument, the default value for this function is `na.rm = TRUE`.

```
> cor(parenthood2, use = "complete.obs")
      dan.sleep baby.sleep  dan.grump       day
dan.sleep   1.00000000  0.6394985 -0.89951468  0.06132891
baby.sleep  0.63949845  1.0000000 -0.58656066  0.14555814
dan.grump  -0.89951468 -0.5865607  1.00000000 -0.06816586
day        0.06132891  0.1455581 -0.06816586  1.00000000
```

The other possibility that we care about, and the one that tends to get used more often in practice, is to set `use = "pairwise.complete.obs"`. When we do that, R only looks at the variables that it's trying to correlate when determining what to drop. So, for instance, since the only missing value for observation 1 of `parenthood2` is for `baby.sleep` R will only drop observation 1 when `baby.sleep` is one of the variables involved: and so R keeps observation 1 when trying to correlate `dan.sleep` and `dan.grump`. When we do it this way, here's what we get:

```
> cor(parenthood2, use = "pairwise.complete.obs")
      dan.sleep baby.sleep  dan.grump       day
dan.sleep   1.00000000  0.61472303 -0.903442442 -0.076796665
baby.sleep  0.61472303  1.00000000 -0.567802669  0.058309485
dan.grump  -0.90344244 -0.56780267  1.000000000  0.005833399
day        -0.07679667  0.05830949  0.005833399  1.000000000
```

Similar, but not quite the same.

The two approaches have different strengths and weaknesses. The “pairwise complete” approach has the advantage that it keeps more observations, so you’re making use of more of your data and (as we’ll discuss in tedious detail in Chapter 10) and it improves the reliability of your estimated correlation. On the other hand, it means that every correlation in your correlation matrix is being computed from a slightly different set of observations, which can be awkward when you want to compare the different correlations that you’ve got.

So which method should you use? It depends a lot on *why* you think your values are missing, and probably depends a little on how paranoid you are. For instance, if you think that the missing values were “chosen” completely randomly²³ then you’ll probably want to use the pairwise method. If you think that missing data are a cue to thinking that the whole observation might be rubbish (e.g., someone just selecting arbitrary responses in your questionnaire), but that there’s no pattern to which observations are “rubbish” then it’s probably safer to keep only those observations that are complete. If you think there’s something systematic going on, in that some observations are more likely to be missing than others, then you have a much trickier problem to solve, and one that is beyond the scope of this book.

5.9 _____

Summary

- Measures of central tendency: mean, median and mode. (Section 5.1)
- Measures of variability: range, standard deviation, interquartile reange (Section 5.2)
- Getting summaries of variables in R (Section 5.4 and 5.5)
- Standard scores (Section 5.6)
- Correlations and rank order correlations (Section 5.7)

²³The technical term here is “missing completely at random” (often written MCAR for short). Makes sense, I suppose, but it does sound ungrammatical to me.

- Basics of handling missing data (Section 5.8)

6. Drawing graphs

Visualising data is one of the most important tasks facing the data analyst. It's important for two distinct but closely related reasons. Firstly, there's the matter of drawing "presentation graphics": displaying your data in a clean, visually appealing fashion makes it easier for your reader to understand what you're trying to tell them. Equally important, perhaps even more important, is the fact that drawing graphs helps *you* to understand the data. To that end, it's important to draw "exploratory graphics" that help you learn about the data as you go about analysing it. These points might seem pretty obvious, but I cannot count the number of times I've seen people forget them.

To give a sense of the importance of this chapter, I want to start with a classic illustration of just how powerful a good graph can be. To that end, Figure 6.1 shows a redrawing of one of the most famous data visualisations of all time: John Snow's 1854 map of cholera deaths. The map is elegant in its simplicity. In the background we have a street map, which helps orient the viewer. Over the top, we see a large number of small dots, each one representing the location of a cholera case. The larger symbols show the location of water pumps, labelled by name. Even the most casual inspection of the graph makes it very clear that the source of the outbreak is almost certainly the Broad Street pump. Upon viewing this graph, Dr Snow arranged to have the handle removed from the pump, ending the outbreak that had killed over 500 people. Such is the power of a good data visualisation.

The goals in this chapter are twofold: firstly, to discuss several fairly standard graphs that we use a lot when analysing and presenting data, and secondly, to show you how to create these graphs in R. The graphs themselves tend to be pretty straightforward, so in that respect this chapter is pretty simple. Where people usually struggle is learning how to produce graphs, and especially, learning how to produce good graphs.¹ Fortunately, learning how to draw graphs in R is reasonably simple, as long as you're not too picky about what your graph looks like. What I mean when I say this is that R has a lot of *very* good graphing functions, and most of the time you can produce a clean, high-quality graphic without having to learn very much about the low-level details of how R handles graphics. Unfortunately, on those occasions when you do want to do something non-standard, or if you need to make highly specific changes to the figure, you actually do need to learn a fair bit about the these details; and those details are both complicated and boring. With that in mind, the structure of this chapter is as follows: I'll start out by giving you a very quick overview of how graphics work in R. I'll then discuss several different kinds of graph and how to draw them, as well as showing the basics of how to customise these plots. I'll then talk in more detail about R graphics, discussing some of those complicated and boring issues. In a future version of this book, I intend to finish this chapter off by talking about what makes a good or a bad graph, but I haven't yet had the time to write that section.

¹I should add that this isn't unique to R. Like everything in R there's a pretty steep learning curve to learning how to draw graphs, and like always there's a massive payoff at the end in terms of the quality of what you can produce. But to be honest, I've seen the same problems show up regardless of what system people use. I suspect that the hardest thing to do is to force yourself to take the time to think deeply about what your graphs are doing. I say that in full knowledge that only about half of my graphs turn out as well as they ought to. Understanding what makes a good graph is easy: actually designing a good graph is *hard*.

Snow's cholera map of London

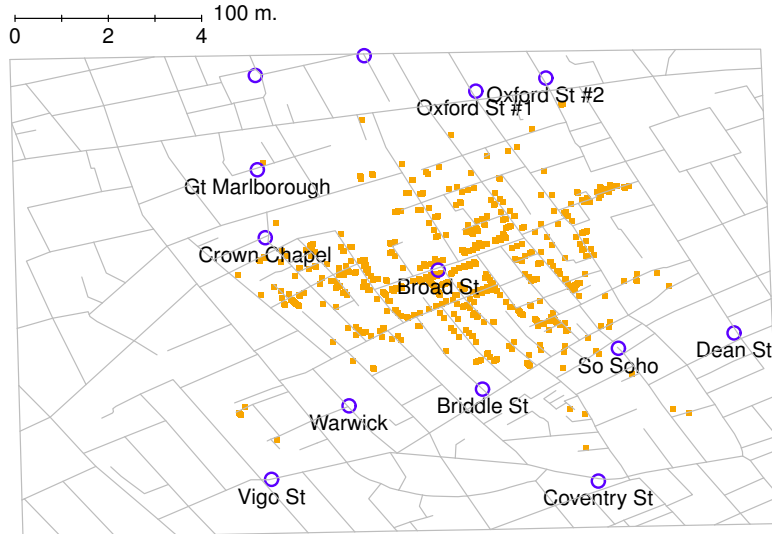


Figure 6.1: A stylised redrawing of John Snow's original cholera map. Each small dot represents the location of a cholera case, and each large circle shows the location of a well. As the plot makes clear, the cholera outbreak is centred very closely on the Broad St pump. This image uses the data from the `HistData` package (Friendly, 2011), and was drawn using minor alterations to the commands provided in the help files. Note that Snow's original hand drawn map used different symbols and labels, but you get the idea.

.....

6.1

An overview of R graphics

Reduced to its simplest form, you can think of an R graphic as being much like a painting. You start out with an empty canvas. Every time you use a graphics function, it paints some new things onto your canvas. Later on, you can paint more things over the top if you want; but just like painting, you can't "undo" your strokes. If you make a mistake, you have to throw away your painting and start over. Fortunately, this is way more easy to do when using R than it is when painting a picture in real life: you delete the plot and then type a new set of commands.² This way of thinking about drawing graphs is referred to as the **painter's model**. So far, this probably doesn't sound particularly complicated, and for the vast majority of graphs you'll want to draw it's exactly as simple as it sounds. Much like painting in real life, the headaches usually start when we dig into details. To see why, I'll expand this "painting metaphor" a bit further just to show you the basics of what's going on under the hood, but before I do I want to stress that you really don't need to understand all these complexities in order to draw graphs.

²Or, since you can always use the up and down keys to scroll through your recent command history, you can just pull up your most recent commands and edit them to fix your mistake. It becomes even easier once you start using scripts (Section 8.1), since all you have to do is edit your script and then run it again.

I'd been using R for years before I even realised that most of these issues existed! However, I don't want you to go through the same pain I went through every time I inadvertently discovered one of these things, so here's a quick overview.

Firstly, if you want to paint a picture, you need to paint it **on** something. In real life, you can paint on lots of different things. Painting onto canvas isn't the same as painting onto paper, and neither one is the same as painting on a wall. In R, the thing that you paint your graphic onto is called a **device**. For most applications that we'll look at in this book, this "device" will be a window on your computer. If you're using Windows as your operating system, then the name for this device is `windows`; on a Mac it's called `quartz` because that's the name of the software that the Mac OS uses to draw pretty pictures; and on Linux/Unix, you're probably using `X11`. On the other hand, if you're using Rstudio (regardless of which operating system you're on), there's a separate device called `RStudioGD` that forces R to paint inside the "plots" panel in Rstudio. However, from the computers perspective there's nothing terribly special about drawing pictures on screen: and so R is quite happy to paint pictures directly into a file. R can paint several different types of image files: `jpeg`, `png`, `pdf`, `postscript`, `tiff` and `bmp` files are all among the options that you have available to you. For the most part, these different devices all behave the same way, so you don't really need to know much about the differences between them when learning how to draw pictures. But, just like real life painting, sometimes the specifics do matter. Unless stated otherwise, you can assume that I'm drawing a picture on screen, using the appropriate device (i.e., `windows`, `quartz`, `X11` or `RStudioGD`). One the rare occasions where these behave differently from one another, I'll try to point it out in the text.

Secondly, when you paint a picture you need to paint it **with** something. Maybe you want to do an oil painting, but maybe you want to use watercolour. And, generally speaking, you pretty much have to pick one or the other. The analog to this in R is a "graphics system". A graphics system defines a collection of very **low-level graphics** commands about what to draw and where to draw it. Something that surprises most new R users is the discovery that R actually has *two* completely independent graphics systems, known as **traditional graphics** (in the `graphics` package) and **grid graphics** (in the `grid` package).³ Not surprisingly, the traditional graphics system is the older of the two: in fact, it's actually older than R since it has its origins in S, the system from which R is descended. Grid graphics are newer, and in some respects more powerful, so many of the more recent, fancier graphical tools in R make use of grid graphics. However, grid graphics are somewhat more complicated beasts, so most people start out by learning the traditional graphics system. Nevertheless, as long as you don't want to use any low-level commands yourself, then you don't really need to care about whether you're using traditional graphics or grid graphics. However, the moment you do want to tweak your figure by using some low-level commands you do need to care. Because these two different systems are pretty much incompatible with each other, there's a pretty big divide in R graphics universe. Unless stated otherwise, you can assume that everything I'm saying pertains to traditional graphics.

Thirdly, a painting is usually done in a particular **style**. Maybe it's a still life, maybe it's an impressionist piece, or maybe you're trying to annoy me by pretending that cubism is a legitimate artistic style. Regardless, each artistic style imposes some overarching aesthetic and perhaps even constraints on what can (or should) be painted using that style. In the same vein, R has quite a number of different packages, each of which provide a collection of **high-level graphics** commands. A single high-level command is capable of drawing an entire graph, complete with a range of customisation options. Most but not all of the high-level commands that I'll talk about in this book come from the `graphics` package itself, and so belong to the world of traditional graphics. These commands all tend to share a common visual style, although there are a few graphics that I'll use that come from other packages that differ in style somewhat. On the other side of the great divide, the grid universe relies heavily on two different packages – `lattice` and `ggplot2` – each of which provides a quite different visual style. As you've probably guessed,

³Of course, even that is a slightly misleading description, since some R graphics tools make use of external graphical rendering systems like OpenGL (e.g., the `rgl` package). I absolutely will not be talking about OpenGL or the like in this book, but as it happens there is one graph in this book that relies on them: Figure 15.4.

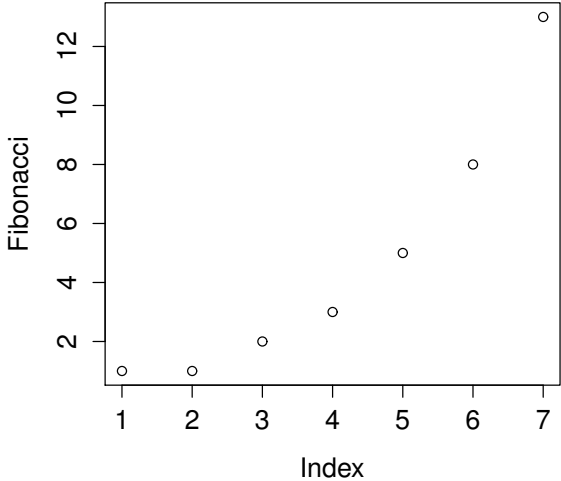


Figure 6.2: Our first plot

there's a whole separate bunch of functions that you'd need to learn if you want to use `lattice` graphics or make use of the `gplots2`. However, for the purposes of this book I'll restrict myself to talking about the basic `graphics` tools.

At this point, I think we've covered more than enough background material. The point that I'm trying to make by providing this discussion isn't to scare you with all these horrible details, but rather to try to convey to you the fact that R doesn't really provide a single coherent graphics system. Instead, R itself provides a platform, and different people have built different graphical tools using that platform. As a consequence of this fact, there's two different universes of graphics, and a great multitude of packages that live in them. At this stage you don't need to understand these complexities, but it's useful to know that they're there. But for now, I think we can be happy with a simpler view of things: we'll draw pictures on screen using the traditional graphics system, and as much as possible we'll stick to high level commands only.

So let's start painting.

6.2

An introduction to plotting

Before I discuss any specialised graphics, let's start by drawing a few very simple graphs just to get a feel for what it's like to draw pictures using R. To that end, let's create a small vector `Fibonacci` that contains a few numbers we'd like R to draw for us. Then, we'll ask R to `plot()` those numbers:

```
> Fibonacci <- c( 1,1,2,3,5,8,13 )
> plot( Fibonacci )
```

The result is Figure 6.2. As you can see, what R has done is plot the *values* stored in the `Fibonacci` variable on the vertical axis (y-axis) and the corresponding *index* on the horizontal axis (x-axis). In other words, since the 4th element of the vector has a value of 3, we get a dot plotted at the location (4,3). That's pretty straightforward, and the image in Figure 6.2 is probably pretty close to what you would have had in mind when I suggested that we plot the `Fibonacci` data. However, there's quite a lot of customisation options available to you, so we should probably spend a bit of time looking at some of those options. So, be warned: this ends up being a fairly long section, because there's so many possibilities open to you. Don't let it overwhelm you though... while all of the options discussed here are handy to know about, you can get by just fine only knowing a few of them. The only reason I've included all this stuff right at the beginning is that it ends up making the rest of the chapter a lot more readable!

6.2.1 A tedious digression

Before we go into any discussion of customising plots, we need a little more background. The important thing to note when using the `plot()` function, is that it's another example of a *generic* function (Section 4.11), much like `print()` and `summary()`, and so its behaviour changes depending on what kind of input you give it. However, the `plot()` function is somewhat fancier than the other two, and its behaviour depends on *two* arguments, `x` (the first input, which is required) and `y` (which is optional). This makes it (a) extremely powerful once you get the hang of it, and (b) hilariously unpredictable, when you're not sure what you're doing. As much as possible, I'll try to make clear what type of inputs produce what kinds of outputs. For now, however, it's enough to note that I'm only doing very basic plotting, and as a consequence all of the work is being done by the `plot.default()` function.

What kinds of customisations might we be interested in? If you look at the help documentation for the default plotting method (i.e., type `?plot.default` or `help("plot.default")`) you'll see a very long list of arguments that you can specify to customise your plot. I'll talk about several of them in a moment, but first I want to point out something that might seem quite wacky. When you look at all the different options that the help file talks about, you'll notice that *some* of the options that it refers to are "proper" arguments to the `plot.default()` function, but it also goes on to mention a bunch of things that *look* like they're supposed to be arguments, but they're not listed in the "Usage" section of the file, and the documentation calls them **graphical parameters** instead. Even so, it's usually possible to treat them as if they were arguments of the plotting function. Very odd. In order to stop my readers trying to find a brick and look up my home address, I'd better explain what's going on; or at least give the basic gist behind it.

What exactly is a graphical parameter? Basically, the idea is that there are some characteristics of a plot which are pretty universal: for instance, regardless of what kind of graph you're drawing, you probably need to specify what colour to use for the plot, right? So you'd expect there to be something like a `col` argument to every single graphics function in R? Well, sort of. In order to avoid having hundreds of arguments for every single function, what R does is refer to a bunch of these "graphical parameters" which are pretty general purpose. Graphical parameters can be changed directly by using the low-level `par()` function, which I discuss briefly in Section 6.7.1 though not in a lot of detail. If you look at the help files for graphical parameters (i.e., type `?par`) you'll see that there's *lots* of them. Fortunately, (a) the default settings are generally pretty good so you can ignore the majority of the parameters, and (b) as you'll see as we go through this chapter, you very rarely need to use `par()` directly, because you can "pretend" that graphical parameters are just additional arguments to your high-level function (e.g. `plot.default()`). In short... yes, R does have these wacky "graphical parameters" which can be quite confusing. But in most basic uses of the plotting functions, you can act as if they were just undocumented additional arguments

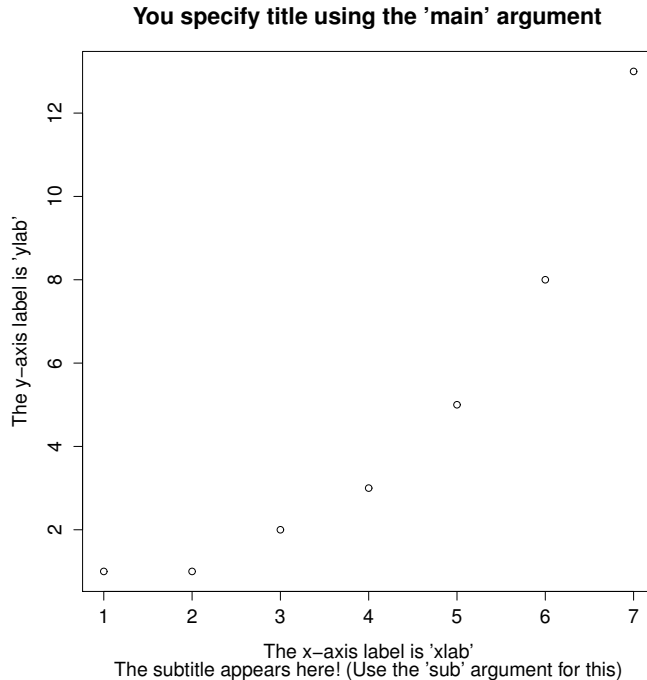


Figure 6.3: How to add your own title, subtitle, x-axis label and y-axis label to the plot. I've drawn this figure a bit larger than the last one so that everything fits. Generally, you don't want your titles and subtitle to extend beyond the range of the actual plot. The results aren't pretty when that happens.

to your function.

6.2.2 Customising the title and the axis labels

One of the first things that you'll find yourself wanting to do when customising your plot is to label it better. You might want to specify more appropriate axis labels, add a title or add a subtitle. The arguments that you need to specify to make this happen are:

- **main**. A character string containing the title.
- **sub**. A character string containing the subtitle.
- **xlab**. A character string containing the x-axis label.
- **ylab**. A character string containing the y-axis label.

These aren't graphical parameters, they're arguments to the high-level function. However, because the high-level functions all rely on the same low-level function to do the drawing⁴ the names of these arguments are identical for pretty much every high-level function I've come across. Let's have a look at what happens when we make use of all these arguments. Here's the command...

⁴The low-level function that does this is called `title()` in case you ever need to know, and you can type `?title` to find out a bit more detail about what these arguments do.

```

> plot( x = Fibonacci,
+       main = "You specify title using the 'main' argument",
+       sub = "The subtitle appears here! (Use the 'sub' argument for this)",
+       xlab = "The x-axis label is 'xlab'",
+       ylab = "The y-axis label is 'ylab'"
+ )

```

The picture that this draws is shown in Figure 6.3. It's more or less as you'd expect. The plot itself is identical to the one we drew in Figure 6.2, except for the fact that we've changed the axis labels, and added a title and a subtitle. Even so, there's a couple of interesting features worth calling your attention to. Firstly, notice that the subtitle is drawn below the plot, which I personally find annoying; as a consequence I almost never use subtitles. You may have a different opinion, of course, but the important thing is that you remember where the subtitle actually goes. Secondly, notice that R has decided to use boldface text and a larger font size for the title. This is one of my most hated default settings in R graphics, since I feel that it draws too much attention to the title. Generally, while I do want my reader to look at the title, I find that the R defaults are a bit overpowering, so I often like to change the settings. To that end, there are a bunch of graphical parameters that you can use to customise the font style:

- *Font styles*: `font.main`, `font.sub`, `font.lab`, `font.axis`. These four parameters control the font style used for the plot title (`font.main`), the subtitle (`font.sub`), the axis labels (`font.lab`: note that you can't specify separate styles for the x-axis and y-axis without using low level commands), and the numbers next to the tick marks on the axis (`font.axis`). Somewhat irritatingly, these arguments are numbers instead of meaningful names: a value of 1 corresponds to plain text, 2 means boldface, 3 means italic and 4 means bold italic.
- *Font colours*: `col.main`, `col.sub`, `col.lab`, `col.axis`. These parameters do pretty much what the name says: each one specifies a colour in which to type each of the different bits of text. Conveniently, R has a very large number of named colours (type `colours()` to see a list of over 650 colour names that R knows), so you can use the English language name of the colour to select it.⁵ Thus, the parameter value here string like "`red`", "`gray25`" or "`springgreen4`" (yes, R really does recognise four different shades of "spring green").
- *Font size*: `cex.main`, `cex.sub`, `cex.lab`, `cex.axis`. Font size is handled in a slightly curious way in R. The "cex" part here is short for "character expansion", and it's essentially a magnification value. By default, all of these are set to a value of 1, except for the font title: `cex.main` has a default magnification of 1.2, which is why the title font is 20% bigger than the others.
- *Font family*: `family`. This argument specifies a font family to use: the simplest way to use it is to set it to "`sans`", "`serif`", or "`mono`", corresponding to a san serif font, a serif font, or a monospaced font. If you want to, you can give the name of a specific font, but keep in mind that different operating systems use different fonts, so it's probably safest to keep it simple. Better yet, unless you have some deep objections to the R defaults, just ignore this parameter entirely. That's what I usually do.

To give you a sense of how you can use these parameters to customise your titles, the following command can be used to draw Figure 6.4:

```

> plot( x = Fibonacci,                                # the data to plot
+       main = "The first 7 Fibonacci numbers",      # the title
+       xlab = "Position in the sequence",            # x-axis label
+       ylab = "The Fibonacci number",                # y-axis label

```

⁵On the off chance that this isn't enough freedom for you, you can select a colour directly as a "red, green, blue" specification using the `rgb()` function, or as a "hue, saturation, value" specification using the `hsv()` function.

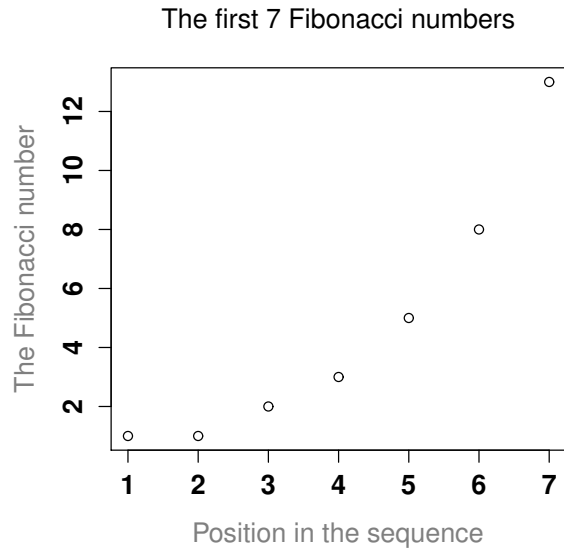


Figure 6.4: How to customise the appearance of the titles and labels.

```
+     font.main = 1,                      # plain text for title
+     cex.main = 1,                      # normal size for title
+     font.axis = 2,                      # bold text for numbering
+     col.lab = "gray50"                 # grey colour for labels
+ )
```

Although this command is quite long, it's not complicated: all it does is override a bunch of the default parameter values. The only difficult aspect to this is that you have to remember what each of these parameters is called, and what all the different values are. And in practice I never remember: I have to look up the help documentation every time, or else look it up in this book.

6.2.3 Changing the plot type

Adding and customising the titles associated with the plot is one way in which you can play around with what your picture looks like. Another thing that you'll want to do is customise the appearance of the actual plot! To start with, let's look at the single most important options that the `plot()` function (or, recalling that we're dealing with a generic function, in this case the `plot.default()` function, since that's the one doing all the work) provides for you to use, which is the `type` argument. The `type` argument specifies the visual style of the plot. The possible values for this are:

- `type = "p"`. Draw the points only.
- `type = "l"`. Draw a line through the points.
- `type = "o"`. Draw the line over the top of the points.
- `type = "b"`. Draw both points and lines, but don't overplot.
- `type = "h"`. Draw “histogram-like” vertical bars.

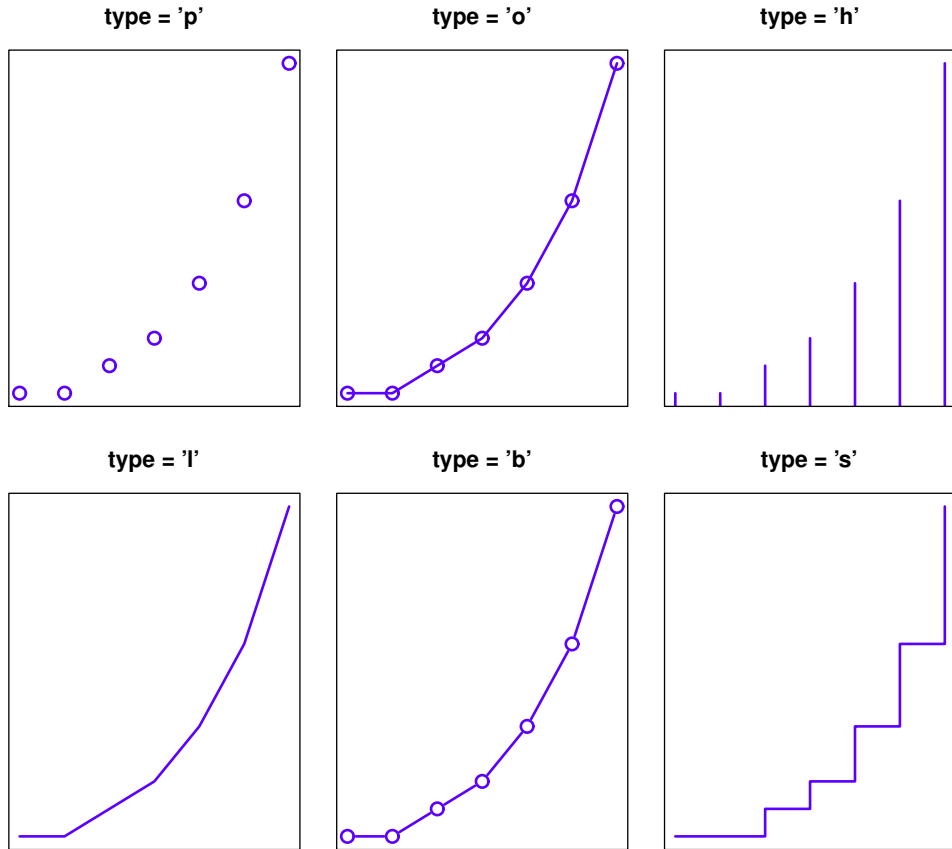


Figure 6.5: Changing the `type` of the plot.

- `type = "s"`. Draw a staircase, going horizontally then vertically.
- `type = "S"`. Draw a Staircase, going vertically then horizontally.
- `type = "c"`. Draw only the connecting lines from the “b” version.
- `type = "n"`. Draw nothing. (Apparently this is useful sometimes?)

The simplest way to illustrate what each of these really looks like is just to draw them. To that end, Figure 6.5 shows the same Fibonacci data, drawn using six different `types` of plot. As you can see, by altering the `type` argument you can get a qualitatively different appearance to your plot. In other words, as far as R is concerned, the only difference between a scatterplot (like the ones we drew in Section 5.7) and a line plot is that you draw a scatterplot by setting `type = "p"` and you draw a line plot by setting `type = "l"`. However, that doesn’t imply that *you* should think of them as being equivalent to each other. As you can see by looking at Figure 6.5, a line plot implies that there is some notion of continuity from one point to the next, whereas a scatterplot does not.

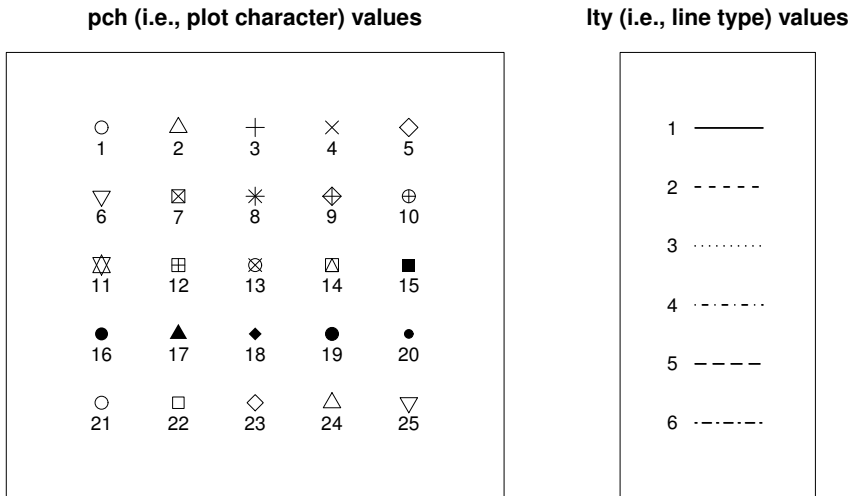


Figure 6.6: Changing the `pch` parameter (panel a) or the `lty` parameter (panel b).

6.2.4 Changing other features of the plot

In Section 6.2.2 we talked about a group of graphical parameters that are related to the formatting of titles, axis labels etc. The second group of parameters I want to discuss are those related to the formatting of the plot itself:

- *Colour of the plot: col*. As we saw with the previous colour-related parameters, the simplest way to specify this parameter is using a character string: e.g., `col = "blue"`. It's a pretty straightforward parameter to specify: the only real subtlety is that every high-level function tends to draw a different “thing” as it’s output, and so this parameter gets interpreted a little differently by different functions. However, for the `plot.default()` function it’s pretty simple: the `col` argument refers to the colour of the points and/or lines that get drawn!
- *Character used to plot points: pch*. The `plot` **character** parameter is a number, usually between 1 and 25. What it does is tell R what symbol to use to draw the points that it plots. The simplest way to illustrate what the different values do is with a picture. Figure 6.6a shows the first 25 plotting characters. The default plotting character is a hollow circle (i.e., `pch = 1`).
- *Plot size: cex*. This parameter describes a `character` expansion factor (i.e., magnification) for the plotted characters. By default `cex=1`, but if you want bigger symbols in your graph you should specify a larger value.
- *Line type: lty*. The `line` **type** parameter describes the kind of line that R draws. It has seven values which you can specify using a number between 0 and 7, or using a meaningful character string: `"blank"`, `"solid"`, `"dashed"`, `"dotted"`, `"dotdash"`, `"longdash"`, or `"twodash"`. Note that the “blank” version (value 0) just means that R doesn’t draw the lines at all. The other six versions are shown in Figure 6.6b.

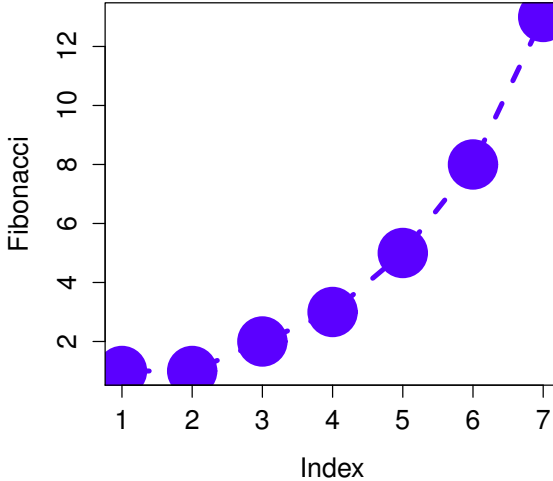


Figure 6.7: Customising various aspects to the plot itself.

- *Line width:* `lwd`. The last graphical parameter in this category that I want to mention is the `line width` parameter, which is just a number specifying the width of the line. The default value is 1. Not surprisingly, larger values produce thicker lines and smaller values produce thinner lines. Try playing around with different values of `lwd` to see what happens.

To illustrate what you can do by altering these parameters, let's try the following command:

```
> plot( x = Fibonacci,    # the data set
+       type = "b",        # plot both points and lines
+       col = "blue",       # change the plot colour to blue
+       pch = 19,          # plotting character is a solid circle
+       cex = 5,           # plot it at 5x the usual size
+       lty = 2,           # change line type to dashed
+       lwd = 4            # change line width to 4x the usual
+ )
```

The output is shown in Figure 6.7.

6.2.5 Changing the appearance of the axes

There are several other possibilities worth discussing. Ignoring graphical parameters for the moment, there's a few other arguments to the `plot.default()` function that you might want to use. As before, many of these are standard arguments that are used by a lot of high level graphics functions:

- *Changing the axis scales:* `xlim`, `ylim`. Generally R does a pretty good job of figuring out where to set the edges of the plot. However, you can override its choices by setting the `xlim` and `ylim`

arguments. For instance, if I decide I want the vertical scale of the plot to run from 0 to 100, then I'd set `ylim = c(0, 100)`.

- *Suppress labelling:* `ann`. This is a logical-valued argument that you can use if you don't want R to include any text for a title, subtitle or axis label. To do so, set `ann = FALSE`. This will stop R from including any text that would normally appear in those places. Note that this will override any of your manual titles. For example, if you try to add a title using the `main` argument, but you also specify `ann = FALSE`, no title will appear.
- *Suppress axis drawing:* `axes`. Again, this is a logical valued argument. Suppose you don't want R to draw any axes at all. To suppress the axes, all you have to do is add `axes = FALSE`. This will remove the axes and the numbering, but not the axis labels (i.e. the `xlab` and `ylab` text). Note that you can get finer grain control over this by specifying the `xaxt` and `yaxt` graphical parameters instead (see below).
- *Include a framing box:* `frame.plot`. Suppose you've removed the axes by setting `axes = FALSE`, but you still want to have a simple box drawn around the plot; that is, you only wanted to get rid of the numbering and the tick marks, but you want to keep the box. To do that, you set `frame.plot = TRUE`.

Note that this list isn't exhaustive. There are a few other arguments to the `plot.default` function that you can play with if you want to, but those are the ones you are probably most likely to want to use. As always, however, if these aren't enough options for you, there's also a number of other graphical parameters that you might want to play with as well. That's the focus of the next section. In the meantime, here's a command that makes use of all these different options:

```
> plot( x = Fibonacci,      # the data
+       xlim = c(0, 15),    # expand the x-scale
+       ylim = c(0, 15),    # expand the y-scale
+       ann = FALSE,        # delete all annotations
+       axes = FALSE,        # delete the axes
+       frame.plot = TRUE   # but include a framing box
+ )
```

The output is shown in Figure 6.8, and it's pretty much exactly as you'd expect. The axis scales on both the horizontal and vertical dimensions have been expanded, the axes have been suppressed as have the annotations, but I've kept a box around the plot.

Before moving on, I should point out that there are several graphical parameters relating to the axes, the box, and the general appearance of the plot which allow finer grain control over the appearance of the axes and the annotations.

- *Suppressing the axes individually:* `xaxt`, `yaxt`. These graphical parameters are basically just fancier versions of the `axes` argument we discussed earlier. If you want to stop R from drawing the vertical axis but you'd like it to keep the horizontal axis, set `yaxt = "n"`. I trust that you can figure out how to keep the vertical axis and suppress the horizontal one!
- *Box type:* `bty`. In the same way that `xaxt`, `yaxt` are just fancy versions of `axes`, the `box` type parameter is really just a fancier version of the `frame.plot` argument, allowing you to specify exactly which out of the four borders you want to keep. The way we specify this parameter is a bit stupid, in my opinion: the possible values are `"o"` (the default), `"l"`, `"7"`, `"c"`, `"u"`, or "]", each of which will draw only those edges that the corresponding character suggests. That is, the letter `"c"` has a top, a bottom and a left, but is blank on the right hand side, whereas `"7"` has a top and a right, but is blank on the left and the bottom. Alternatively a value of `"n"` means that no box will be drawn.

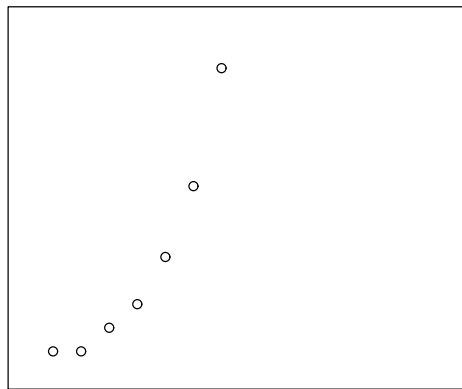


Figure 6.8: Altering the scale and appearance of the plot axes.

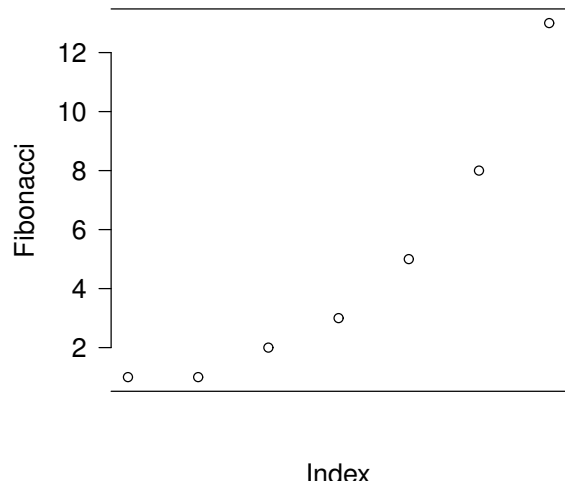


Figure 6.9: Other ways to customise the axes.

- *Orientation of the axis labels `las`.* I presume that the name of this parameter is an acronym of label style or something along those lines; but what it actually does is govern the orientation of the text used to label the individual tick marks (i.e., the numbering, not the `xlab` and `ylab` axis labels). There are four possible values for `las`: A value of 0 means that the labels of both axes are printed parallel to the axis itself (the default). A value of 1 means that the text is always horizontal. A value of 2 means that the labelling text is printed at right angles to the axis. Finally, a value of 3 means that the text is always vertical.

Again, these aren't the only possibilities. There are a few other graphical parameters that I haven't mentioned that you could use to customise the appearance of the axes,⁶ but that's probably enough (or more than enough) for now. To give a sense of how you could use these parameters, let's try the following command:

```
> plot( x = Fibonacci,    # the data
+       xaxt = "n",      # don't draw the x-axis
+       bty = "]",        # keep bottom, right and top of box only
+       las = 1           # rotate the text
+ )
```

The output is shown in Figure 6.9. As you can see, this isn't a very useful plot at all. However, it does illustrate the graphical parameters we're talking about, so I suppose it serves its purpose.

6.2.6 Don't panic

At this point, a lot of readers will be probably be thinking something along the lines of, “if there’s this much detail just for drawing a simple plot, how horrible is it going to get when we start looking at more complicated things?” Perhaps, contrary to my earlier pleas for mercy, you’ve found a brick to hurl and are right now leafing through an Adelaide phone book trying to find my address. Well, fear not! And please, put the brick down. In a lot of ways, we’ve gone through the hardest part: we’ve already covered vast majority of the plot customisations that you might want to do. As you’ll see, each of the other high level plotting commands we’ll talk about will only have a smallish number of additional options. Better yet, even though I’ve told you about a billion different ways of tweaking your plot, you don’t usually need them. So in practice, now that you’ve read over it once to get the gist, the majority of the content of this section is stuff you can safely forget: just remember to come back to this section later on when you want to tweak your plot.

6.3 _____

Histograms

Now that we’ve tamed (or possibly fled from) the beast that is R graphical parameters, let’s talk more seriously about some real life graphics that you’ll want to draw. We begin with the humble **histogram**. Histograms are one of the simplest and most useful ways of visualising data. They make most sense when you have an interval or ratio scale (e.g., the `afl.margins` data from Chapter 5) and what you want to do is get an overall impression of the data. Most of you probably know how histograms work, since they’re so widely used, but for the sake of completeness I’ll describe them. All you do is divide up the possible values into **bins**, and then count the number of observations that fall within each bin. This count is referred to as the frequency of the bin, and is displayed as a bar: in the AFL winning margins data, there are 33 games in which the winning margin was less than 10 points, and it is this fact that is represented

⁶Also, there’s a low level function called `axis()` that allows a lot more control over the appearance of the axes.

by the height of the leftmost bar in Figure 6.10a. Drawing this histogram in R is pretty straightforward. The function you need to use is called `hist()`, and it has pretty reasonable default settings. In fact, Figure 6.10a is exactly what you get if you just type this:

```
> hist( afl.margins ) # panel a
```

Although this image would need a lot of cleaning up in order to make a good presentation graphic (i.e., one you'd include in a report), it nevertheless does a pretty good job of describing the data. In fact, the big strength of a histogram is that (properly used) it does show the entire spread of the data, so you can get a pretty good sense about what it looks like. The downside to histograms is that they aren't very compact: unlike some of the other plots I'll talk about it's hard to cram 20-30 histograms into a single image without overwhelming the viewer. And of course, if your data are nominal scale (e.g., the `afl.finalists` data) then histograms are useless.

The main subtlety that you need to be aware of when drawing histograms is determining where the `breaks` that separate bins should be located, and (relatedly) how many breaks there should be. In Figure 6.10a, you can see that R has made pretty sensible choices all by itself: the breaks are located at 0, 10, 20, ... 120, which is exactly what I would have done had I been forced to make a choice myself. On the other hand, consider the two histograms in Figure 6.10b and 6.10c, which I produced using the following two commands:

```
> hist( x = afl.margins, breaks = 3 )      # panel b
> hist( x = afl.margins, breaks = 0:116 ) # panel c
```

In Figure 6.10c, the bins are only 1 point wide. As a result, although the plot is very informative (it displays the entire data set with no loss of information at all!) the plot is very hard to interpret, and feels quite cluttered. On the other hand, the plot in Figure 6.10b has a bin width of 50 points, and has the opposite problem: it's very easy to "read" this plot, but it doesn't convey a lot of information. One gets the sense that this histogram is hiding too much. In short, the way in which you specify the breaks has a big effect on what the histogram looks like, so it's important to make sure you choose the breaks sensibly. In general R does a pretty good job of selecting the breaks on its own, since it makes use of some quite clever tricks that statisticians have devised for automatically selecting the right bins for a histogram, but nevertheless it's usually a good idea to play around with the breaks a bit to see what happens.

There is one fairly important thing to add regarding how the `breaks` argument works. There are two different ways you can specify the breaks. You can either specify *how many* breaks you want (which is what I did for panel b when I typed `breaks = 3`) and let R figure out where they should go, or you can provide a vector that tells R exactly where the breaks should be placed (which is what I did for panel c when I typed `breaks = 0:116`). The behaviour of the `hist()` function is slightly different depending on which version you use. If all you do is tell it *how many* breaks you want, R treats it as a "suggestion" not as a demand. It assumes you want "approximately 3" breaks, but if it doesn't think that this would look very pretty on screen, it picks a different (but similar) number. It does this for a sensible reason – it tries to make sure that the breaks are located at sensible values (like 10) rather than stupid ones (like 7.224414). And most of the time R is right: usually, when a human researcher says "give me 3 breaks", he or she really does mean "give me approximately 3 breaks, and don't put them in stupid places". However, sometimes R is dead wrong. Sometimes you really do mean "exactly 3 breaks", and you know precisely where you want them to go. So you need to invoke "real person privilege", and order R to do what it's bloody well told. In order to do that, you *have* to input the full vector that tells R exactly where you want the breaks. If you do that, R will go back to behaving like the nice little obedient calculator that it's supposed to be.

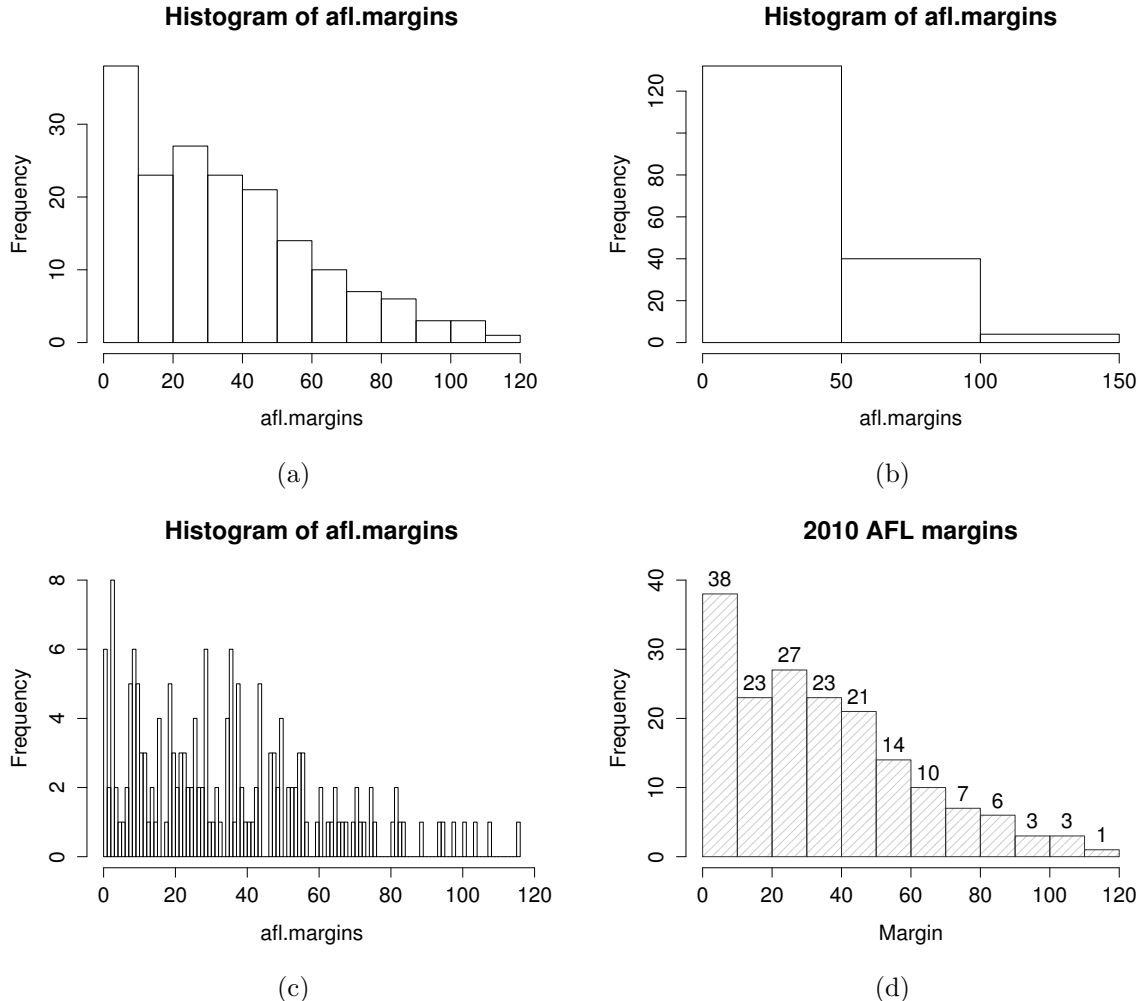


Figure 6.10: Four different histograms of the `afl.margins` variable: (a) the default histogram that R produces, (b) a histogram with too few bins, (c) a histogram with too many bins, and (d) a “prettier” histogram making use of various optional arguments to `hist()`. Overall I think that panel d makes a pretty good picture, though if I were to start being picky I wouldn’t have the title in boldface text (the eye is drawn to the title and not to the picture), and I’d try to make the labels above the bars a little less prominent as well.

6.3.1 Visual style of your histogram

Okay, so at this point we can draw a basic histogram, and we can alter the number and even the location of the `breaks`. However, the visual style of the histograms shown in Figure 6.10a-c could stand to be improved. We can fix this by making use of some of the other arguments to the `hist()` function. Most of the things you might want to try doing have already been covered in Section 6.2, but there's a few new things:

- *Shading lines:* `density`, `angle`. You can add diagonal lines to shade the bars: the `density` value is a number indicating how many lines per inch R should draw (the default value of `NULL` means no lines), and the `angle` is a number indicating how many degrees from horizontal the lines should be drawn at (default is `angle = 45` degrees).
- *Specifics regarding colours:* `col`, `border`. You can also change the colours: in this instance the `col` parameter sets the colour of the shading (either the shading lines if there are any, or else the colour of the interior of the bars if there are not), and the `border` argument sets the colour of the edges of the bars.
- *Labelling the bars:* `labels`. You can also attach labels to each of the bars using the `labels` argument. The simplest way to do this is to set `labels = TRUE`, in which case R will add a number just above each bar, that number being the exact number of observations in the bin. Alternatively, you can choose the labels yourself, by inputting a vector of strings, e.g., `labels = c("label 1", "label 2", "etc")`

Not surprisingly, this doesn't exhaust the possibilities. If you type `help("hist")` or `?hist` and have a look at the help documentation for histograms, you'll see a few more options. A histogram that makes use of the histogram-specific customisations as well as several of the options we discussed in Section 6.2 is shown in Figure 6.10d. The R command that I used to draw it is this:

```
> hist( x = afl.margins,
+       main = "2010 AFL margins", # title of the plot
+       xlab = "Margin",          # set the x-axis label
+       density = 10,            # draw shading lines: 10 per inch
+       angle = 40,              # set the angle of the shading lines is 40 degrees
+       border = "gray20",        # set the colour of the borders of the bars
+       col = "gray80",           # set the colour of the shading lines
+       labels = TRUE,            # add frequency labels to each bar
+       ylim = c(0,40)            # change the scale of the y-axis
+ )
```

Overall, this is a much nicer histogram than the default ones.

6.4

Stem and leaf plots

Histograms are one of the most widely used methods for displaying the observed values for a variable. They're simple, pretty, and very informative. However, they do take a little bit of effort to draw. Sometimes it can be quite useful to make use of simpler, if less visually appealing, options. One such alternative is the **stem and leaf plot**. To a first approximation you can think of a stem and leaf plot as a kind of text-based histogram. Stem and leaf plots aren't used as widely these days as they were 30 years ago, since it's now just as easy to draw a histogram as it is to draw a stem and leaf plot. Not only

that, they don't work very well for larger data sets. As a consequence you probably won't have as much of a need to use them yourself, though you may run into them in older publications. These days, the only real world situation where I use them is if I have a small data set with 20-30 data points and I don't have a computer handy, because it's pretty easy to quickly sketch a stem and leaf plot by hand.

With all that as background, let's have a look at stem and leaf plots. The AFL margins data contains 176 observations, which is at the upper end for what you can realistically plot this way. The function in R for drawing stem and leaf plots is called `stem()` and if we ask for a stem and leaf plot of the `afl.margins` data, here's what we get:

```
> stem( afl.margins )

The decimal point is 1 digit(s) to the right of the |

 0 | 00111122333333344567788888999999
 1 | 0000011122234456666899999
 2 | 00011222333445566667788999999
 3 | 0122355566666678888899
 4 | 012334444477788899
 5 | 00002233445556667
 6 | 0113455678
 7 | 01123556
 8 | 122349
 9 | 458
10 | 148
11 | 6
```

The values to the left of the `|` are called **stems** and the values to the right are called **leaves**. If you just look at the shape that the leaves make, you can see something that looks a lot like a histogram made out of numbers, just rotated by 90 degrees. But if you know how to read the plot, there's quite a lot of additional information here. In fact, it's also giving you the actual values of *all* of the observations in the data set. To illustrate, let's have a look at the last line in the stem and leaf plot, namely `11 | 6`. Specifically, let's compare this to the largest values of the `afl.margins` data set:

```
> max( afl.margins )
[1] 116
```

Hm... `11 | 6` versus `116`. Obviously the stem and leaf plot is trying to tell us that the largest value in the data set is `116`. Similarly, when we look at the line that reads `10 | 148`, the way we interpret it to note that the stem and leaf plot is telling us that the data set contains observations with values `101`, `104` and `108`. Finally, when we see something like `5 | 00002233445556667` the four `0`s in the the stem and leaf plot are telling us that there are four observations with value `50`.

I won't talk about them in a lot of detail, but I should point out that some customisation options are available for stem and leaf plots in R. The two arguments that you can use to do this are:

- **scale**. Changing the `scale` of the plot (default value is `1`), which is analogous to changing the number of breaks in a histogram. Reducing the scale causes R to reduce the number of stem values (i.e., the number of breaks, if this were a histogram) that the plot uses.
- **width**. The second way that to can customise a stem and leaf plot is to alter the `width` (default value is `80`). Changing the width alters the maximum number of leaf values that can be displayed for any given stem.

However, since stem and leaf plots aren't as important as they used to be, I'll leave it to the interested reader to investigate these options. Try the following two commands to see what happens:

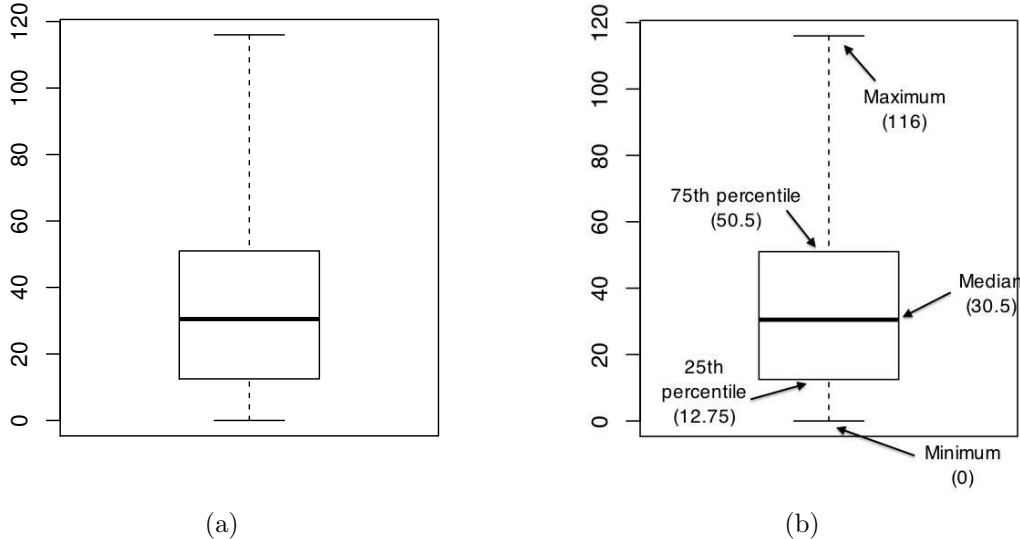


Figure 6.11: A basic boxplot (panel a), plus the same plot with annotations added to explain what aspect of the data set each part of the boxplot corresponds to (panel b).

```
> stem( x = afl.margins, scale = .25 )
> stem( x = afl.margins, width = 20 )
```

The only other thing to note about stem and leaf plots is the line in which R tells you where the decimal point is. If our data set had included only the numbers .11, .15, .23, .35 and .59 and we'd drawn a stem and leaf plot of these data, then R would move the decimal point: the stem values would be 1,2,3,4 and 5, but R would tell you that the decimal point has moved to the left of the | symbol. If you want to see this in action, try the following command:

```
> stem( x = afl.margins / 1000 )
```

The stem and leaf plot itself will look identical to the original one we drew, except for the fact that R will tell you that the decimal point has moved.

6.5 _____

Boxplots

Another alternative to histograms is a **boxplot**, sometimes called a “box and whiskers” plot. Like histograms, they’re most suited to interval or ratio scale data. The idea behind a boxplot is to provide a simple visual depiction of the median, the interquartile range, and the range of the data. And because they do so in a fairly compact way, boxplots have become a very popular statistical graphic, especially during the exploratory stage of data analysis when you’re trying to understand the data yourself. Let’s

have a look at how they work, again using the `afl.margins` data as our example. Firstly, let's actually calculate these numbers ourselves using the `summary()` function:⁷

```
> summary( afl.margins )
   Min. 1st Qu. Median   Mean 3rd Qu.   Max.
   0.00   12.75  30.50  35.30  50.50  116.00
```

So how does a boxplot capture these numbers? The easiest way to describe what a boxplot looks like is just to draw one. The function for doing this in R is (surprise, surprise) `boxplot()`. As always there's a lot of optional arguments that you can specify if you want, but for the most part you can just let R choose the defaults for you. That said, I'm going to override one of the defaults to start with by specifying the `range` option, but for the most part you won't want to do this (I'll explain why in a minute). With that as preamble, let's try the following command:

```
> boxplot( x = afl.margins, range = 100 )
```

What R draws is shown in Figure 6.11a, the most basic boxplot possible. When you look at this plot, this is how you should interpret it: the thick line in the middle of the box is the median; the box itself spans the range from the 25th percentile to the 75th percentile; and the “whiskers” cover the full range from the minimum value to the maximum value. This is summarised in the annotated plot in Figure 6.11b.

In practice, this isn't quite how boxplots usually work. In most applications, the “whiskers” don't cover the full range from minimum to maximum. Instead, they actually go out to the most extreme data point that doesn't exceed a certain bound. By default, this value is 1.5 times the interquartile range, corresponding to a `range` value of 1.5. Any observation whose value falls outside this range is plotted as a circle instead of being covered by the whiskers, and is commonly referred to as an **outlier**. For our AFL margins data, there is one observation (a game with a margin of 116 points) that falls outside this range. As a consequence, the upper whisker is pulled back to the next largest observation (a value of 108), and the observation at 116 is plotted as a circle. This is illustrated in Figure 6.12a. Since the default value is `range = 1.5` we can draw this plot using the simple command

```
> boxplot( afl.margins )
```

6.5.1 Visual style of your boxplot

I'll talk a little more about the relationship between boxplots and outliers in the Section 6.5.2, but before I do let's take the time to clean this figure up. Boxplots in R are extremely customisable. In addition to the usual range of graphical parameters that you can tweak to make the plot look nice, you can also exercise nearly complete control over every element to the plot. Consider the boxplot in Figure 6.12b: in this version of the plot, not only have I added labels (`xlab`, `ylab`) and removed the stupid border (`frame.plot`), I've also dimmed all of the graphical elements of the boxplot except the central bar that plots the median (`border`) so as to draw more attention to the median rather than the rest of the boxplot. You've seen all these options in previous sections in this chapter, so hopefully those customisations won't need any further explanation. However, I've done two new things as well: I've deleted the cross-bars at the top and bottom of the whiskers (known as the “staples” of the plot), and converted the whiskers themselves to solid lines. The arguments that I used to do this are called by the ridiculous names of `staplewex` and `whisklty`,⁸ and I'll explain these in a moment. But first, here's the actual command I used to draw this figure:

⁷R being what it is, it's no great surprise that there's also a `fivenum()` function that does much the same thing.

⁸I realise there's a kind of logic to the way R names are constructed, but they still sound dumb. When I typed this sentence, all I could think was that it sounded like the name of a kids movie if it had been written by Lewis Carroll: “The frabjous gambolles of Staplewex and Whisklty” or something along those lines.

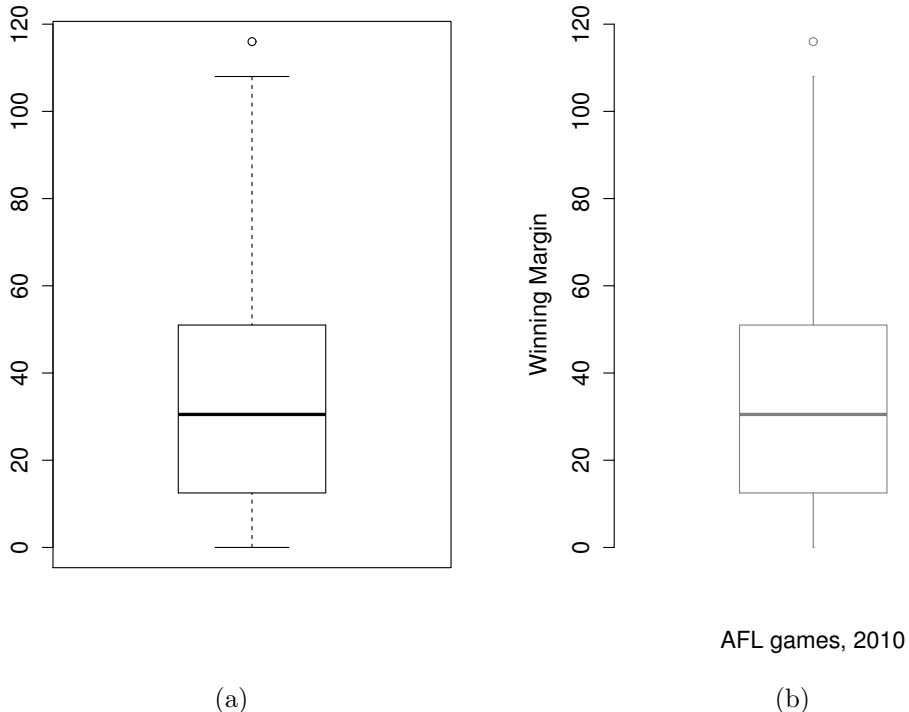


Figure 6.12: By default, R will only extent the whiskers a distance of 1.5 times the interquartile range, and will plot any points that fall outside that range separately (panel a). As we've seen in earlier graphics, the `boxplot()` function lets you customise the plot fairly extensively. This is illustrated in panel b, which shows a much more minimalist boxplot, and attaches informative labels to the graphic.

```
> boxplot( x = afl.margins,           # the data
+          xlab = "AFL games, 2010", # x-axis label
+          ylab = "Winning Margin", # y-axis label
+          border = "grey50",       # dim the border of the box
+          frame.plot = FALSE,     # don't draw a frame
+          staplewex = 0,          # don't draw staples
+          whisklty = 1            # solid line for whisker
+ )
```

Overall, I think the resulting boxplot is a huge improvement in visual design over the default version. In my opinion at least, there's a fairly minimalist aesthetic that governs good statistical graphics. Ideally, every visual element that you add to a plot should convey part of the message. If your plot includes things that don't actually help the reader learn anything new, you should consider removing them. Personally, I can't see the point of the cross-bars on a standard boxplot, so I've deleted them.

Okay, what commands can we use to customise the boxplot? If you type `?boxplot` and flick through the help documentation, you'll notice that it does mention `staplewex` as an argument, but there's no mention of `whisklty`. The reason for this is that the function that handles the drawing is called `bxp()`, so if you type `?bxp` all the gory details appear. Here's the short summary. In order to understand why these arguments have such stupid names, you need to recognise that they're put together from two components.

The first part of the argument name specifies one part of the box plot: `staple` refers to the staples of the plot (i.e., the cross-bars), and `whisk` refers to the whiskers. The second part of the name specifies a graphical parameter: `wex` is a width parameter, and `lty` is a line type parameter. The parts of the plot you can customise are:

- `box`. The box that covers the interquartile range.
- `med`. The line used to show the median.
- `whisk`. The vertical lines used to draw the whiskers.
- `staple`. The cross bars at the ends of the whiskers.
- `out`. The points used to show the outliers.

The actual graphical parameters that you might want to specify are slightly different for each visual element, just because they're different shapes from each other. As a consequence, the following options are available:

- *Width expansion*: `boxwex`, `staplewex`, `outwex`. These are scaling factors that govern the width of various parts of the plot. The default scaling factor is (usually) 0.8 for the box, and 0.5 for the other two. Note that in the case of the outliers this parameter is meaningless unless you decide to draw lines plotting the outliers rather than use points.
- *Line type*: `boxlty`, `medlty`, `whisklty`, `staplelty`, `outlty`. These govern the line type for the relevant elements. The values for this are exactly the same as those used for the regular `lty` parameter, with two exceptions. There's an additional option where you can set `medlty = "blank"` to suppress the median line completely (useful if you want to draw a point for the median rather than plot a line). Similarly, by default the outlier line type is set to `outlty = "blank"`, because the default behaviour is to draw outliers as points instead of lines.
- *Line width*: `boxlwd`, `medlwd`, `whisklwd`, `staplelwd`, `outlwd`. These govern the line widths for the relevant elements, and behave the same way as the regular `lwd` parameter. The only thing to note is that the default value for `medlwd` value is three times the value of the others.
- *Line colour*: `boxcol`, `medcol`, `whiskcol`, `staplecol`, `outcol`. These govern the colour of the lines used to draw the relevant elements. Specify a colour in the same way that you usually do.
- *Fill colour*: `boxfill`. What colour should we use to fill the box?
- *Point character*: `medpch`, `outpch`. These behave like the regular `pch` parameter used to select the plot character. Note that you can set `outpch = NA` to stop R from plotting the outliers at all, and you can also set `medpch = NA` to stop it from drawing a character for the median (this is the default!).
- *Point expansion*: `medcex`, `outcex`. Size parameters for the points used to plot medians and outliers. These are only meaningful if the corresponding points are actually plotted. So for the default boxplot, which includes outlier points but uses a line rather than a point to draw the median, only the `outcex` parameter is meaningful.
- *Background colours*: `medbg`, `outbg`. Again, the background colours are only meaningful if the points are actually plotted.

Taken as a group, these parameters allow you almost complete freedom to select the graphical style for your boxplot that you feel is most appropriate to the data set you're trying to describe. That said, when you're first starting out there's no shame in using the default settings! But if you want to master the art of designing beautiful figures, it helps to try playing around with these parameters to see what works and what doesn't. Finally, I should mention a few other arguments that you might want to make use of:

- `horizontal`. Set this to `TRUE` to display the plot horizontally rather than vertically.

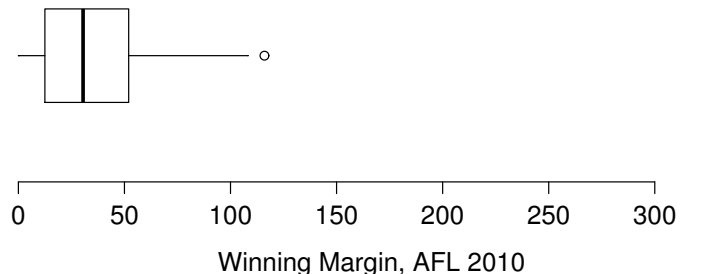


Figure 6.13: A boxplot showing one very suspicious outlier! I've drawn this plot in a similar, minimalist style to the one in Figure 6.12b, but I've used the `horizontal` argument to draw it sideways in order to save space.

- `varwidth`. Set this to `TRUE` to get R to scale the width of each box so that the areas are proportional to the number of observations that contribute to the boxplot. This is only useful if you're drawing multiple boxplots at once (see Section 6.5.3).
- `show.names`. Set this to `TRUE` to get R to attach labels to the boxplots.
- `notch`. If you set `notch = TRUE`, R will draw little notches in the sides of each box. If the notches of two boxplots don't overlap, then there is a "statistically significant" difference between the corresponding medians. If you haven't read Chapter 11, ignore this argument – we haven't discussed statistical significance, so this doesn't mean much to you. I'm mentioning it only because you might want to come back to the topic later on. (see also the `notch.frac` option when you type `?bxp`).

6.5.2 Using box plots to detect outliers

Because the boxplot automatically (unless you change the `range` argument) separates out those observations that lie within a certain range, people often use them as an informal method for detecting **outliers**: observations that are "suspiciously" distant from the rest of the data. Here's an example. Suppose that I'd drawn the boxplot for the AFL margins data, and it came up looking like Figure 6.13. It's pretty clear that something funny is going on with one of the observations. Apparently, there was one game in which the margin was over 300 points! That doesn't sound right to me. Now that I've become suspicious, it's time to look a bit more closely at the data. One function that can be handy for this is the `which()` function; it takes as input a vector of logicals, and outputs the indices of the `TRUE` cases. This is particularly useful in the current context because it lets me do this:

```
> suspicious.cases <- afl.margins > 300
> which( suspicious.cases )
[1] 137
```

although in real life I probably wouldn't bother creating the `suspicious.cases` variable: I'd just cut out the middle man and use a command like `which(afl.margins > 300)`. In any case, what this has done is shown me that the outlier corresponds to game 137. Then, I find the recorded margin for that game:

```
> afl.margins[137]  
[1] 333
```

Hm. That definitely doesn't sound right. So then I go back to the original data source (the internet!) and I discover that the actual margin of that game was 33 points. Now it's pretty clear what happened. Someone must have typed in the wrong number. Easily fixed, just by typing `afl.margins[137] <- 33`. While this might seem like a silly example, I should stress that this kind of thing actually happens a lot. Real world data sets are often riddled with stupid errors, especially when someone had to type something into a computer at some point. In fact, there's actually a name for this phase of data analysis, since in practice it can waste a huge chunk of our time: **data cleaning**. It involves searching for typos, missing data and all sorts of other obnoxious errors in raw data files.⁹

What about the real data? Does the value of 116 constitute a funny observation not? Possibly. As it turns out the game in question was Fremantle v Hawthorn, and was played in round 21 (the second last home and away round of the season). Fremantle had already qualified for the final series and for them the outcome of the game was irrelevant; and the team decided to rest several of their star players. As a consequence, Fremantle went into the game severely underpowered. In contrast, Hawthorn had started the season very poorly but had ended on a massive winning streak, and for them a win could secure a place in the finals. With the game played on Hawthorn's home turf¹⁰ and with so many unusual factors at play, it is perhaps no surprise that Hawthorn annihilated Fremantle by 116 points. Two weeks later, however, the two teams met again in an elimination final on Fremantle's home ground, and Fremantle won comfortably by 30 points.¹¹

So, should we exclude the game from subsequent analyses? If this were a psychology experiment rather than an AFL season, I'd be quite tempted to exclude it because there's pretty strong evidence that Fremantle weren't really trying very hard: and to the extent that my research question is based on an assumption that participants are genuinely trying to do the task. On the other hand, in a lot of studies we're actually interested in seeing the full range of possible behaviour, and that includes situations where people decide not to try very hard: so excluding that observation would be a bad idea. In the context of the AFL data, a similar distinction applies. If I'd been trying to make tips about who would perform well in the finals, I would have (and in fact did) disregard the Round 21 massacre, because it's way too misleading. On the other hand, if my interest is solely in the home and away season itself, I think it would be a shame to throw away information pertaining to one of the most distinctive (if boring) games of the year. In other words, the decision about whether to include outliers or exclude them depends heavily on *why* you think the data look they way they do, and what you want to use the data *for*. Statistical tools can provide an automatic method for suggesting candidates for deletion, but you really need to exercise good judgment here. As I've said before, R is a mindless automaton. It doesn't watch the footy, so it lacks the broader context to make an informed decision. You are *not* a mindless automaton, so you should exercise judgment: if the outlier looks legitimate to you, then keep it. In any case, I'll return to

⁹Sometimes it's convenient to have the boxplot automatically label the outliers for you. The original `boxplot()` function doesn't allow you to do this; however, the `Boxplot()` function in the `car` package does. The design of the `Boxplot()` function is very similar to `boxplot()`. It just adds a few new arguments that allow you to tweak the labelling scheme. I'll leave it to the reader to check this out.

¹⁰Sort of. The game was played in Launceston, which is a de facto home away from home for Hawthorn.

¹¹Contrast this situation with the next largest winning margin in the data set, which was Geelong's 108 point demolition of Richmond in round 6 at their home ground, Kardinia Park. Geelong have been one of the most dominant teams over the last several years, a period during which they strung together an incredible 29-game winning streak at Kardinia Park. Richmond have been useless for several years. This is in no meaningful sense an outlier. Geelong have been winning by these margins (and Richmond losing by them) for quite some time. Frankly I'm surprised that the result wasn't more lopsided: as happened to Melbourne in 2011 when Geelong won by a modest 186 points.

the topic again in Section 15.8, so let's return to our discussion of how to draw boxplots.

6.5.3 Drawing multiple boxplots

One last thing. What if you want to draw multiple boxplots at once? Suppose, for instance, I wanted separate boxplots showing the AFL margins not just for 2010, but for every year between 1987 and 2010. To do that, the first thing we'll have to do is find the data. These are stored in the `aflsmall12.Rdata` file. So let's load it and take a quick peek at what's inside:

```
> load( "aflsmall12.Rdata" )
> who( TRUE )
-- Name --  -- Class --  -- Size --
afl2        data.frame    4296 x 2
$margin     numeric      4296
$year       numeric      4296
```

Notice that `afl2` data frame is pretty big. It contains 4296 games, which is far more than I want to see printed out on my computer screen. To that end, R provides you with a few useful functions to print out only a few of the row in the data frame. The first of these is `head()` which prints out the first 6 rows, of the data frame, like this:

```
> head( afl2 )
  margin year
1     33 1987
2     59 1987
3     45 1987
4     91 1987
5     39 1987
6      1 1987
```

You can also use the `tail()` function to print out the last 6 rows. The `car` package also provides a handy little function called `some()` which prints out a random subset of the rows.

In any case, the important thing is that we have the `afl2` data frame which contains the variables that we're interested in. What we want to do is have R draw boxplots for the `margin` variable, plotted separately for each separate `year`. The way to do this using the `boxplot()` function is to input a `formula` rather than a variable as the input. In this case, the formula we want is `margin ~ year`. So our boxplot command now looks like this:

```
> boxplot( formula = margin ~ year,
+           data = afl2
+ )
```

The result is shown in Figure 6.14.¹² Even this, the default version of the plot, gives a sense of why it's sometimes useful to choose boxplots instead of histograms. Even before taking the time to turn this basic output into something more readable, it's possible to get a good sense of what the data look like from year to year without getting overwhelmed with too much detail. Now imagine what would have happened if I'd tried to cram 24 histograms into this space: no chance at all that the reader is going to learn anything useful.

¹²Actually, there's other ways to do this. If the input argument `x` is a list object (see Section 4.9), the `boxplot()` function will draw a separate boxplot for each variable in that list. Relatedly, since the `plot()` function – which we'll discuss shortly – is a generic (see Section 4.11), you might not be surprised to learn that one of its special cases is a boxplot: specifically, if you use `plot()` where the first argument `x` is a factor and the second argument `y` is numeric, then the result will be a boxplot, showing the values in `y`, with a separate boxplot for each level. For instance, something like `plot(x = afl2$year, y = afl2$margin)` would work.

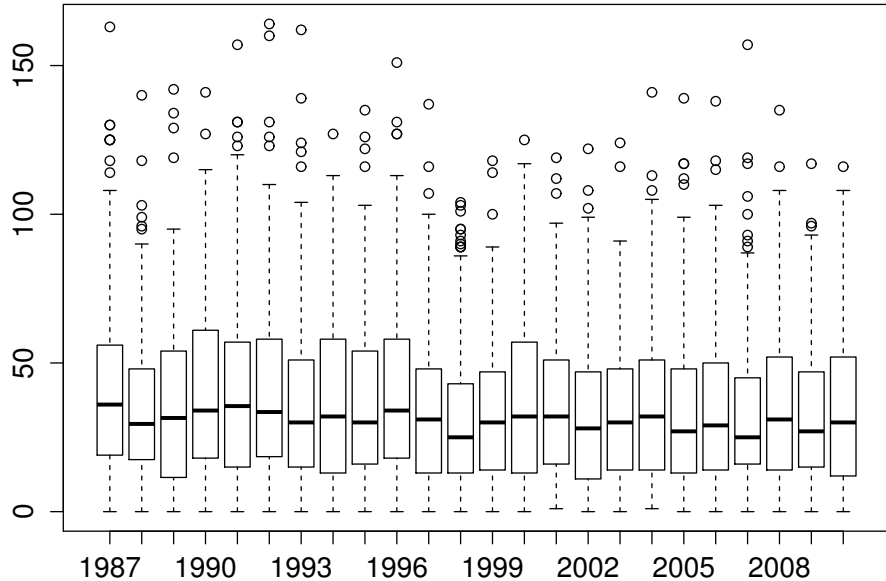


Figure 6.14: Boxplots showing the AFL winning margins for the 24 years from 1987 to 2010 inclusive. This is the default plot created by R, with no annotations added and no changes to the visual design. It's pretty readable, though at a minimum you'd want to include some basic annotations labelling the axes. Compare and contrast with Figure 6.15

That being said, the default boxplot leaves a great deal to be desired in terms of visual clarity. The outliers are too visually prominent, the dotted lines look messy, and the interesting content (i.e., the behaviour of the median and the interquartile range across years) gets a little obscured. Fortunately, this is easy to fix, since we've already covered a lot of tools you can use to customise your output. After playing around with several different versions of the plot, the one I settled on is shown in Figure 6.15. The command I used to produce it is long, but not complicated:

```
> boxplot( formula = margin ~ year,      # the formula
+         data = afl2,                  # the data set
+         xlab = "AFL season",        # x axis label
+         ylab = "Winning Margin",    # y axis label
+         frame.plot = FALSE,        # don't draw a frame
+         staplewex = 0,              # don't draw staples
+         staplecol = "white",       # (fixes a tiny display issue)
+         boxwex = .75,              # narrow the boxes slightly
+         boxfill = "grey80",          # lightly shade the boxes
+         whisklty = 1,               # solid line for whiskers
+         whiskcol = "grey70",        # dim the whiskers
+         boxcol = "grey70",          # dim the box borders
+         outcol = "grey70",          # dim the outliers
```

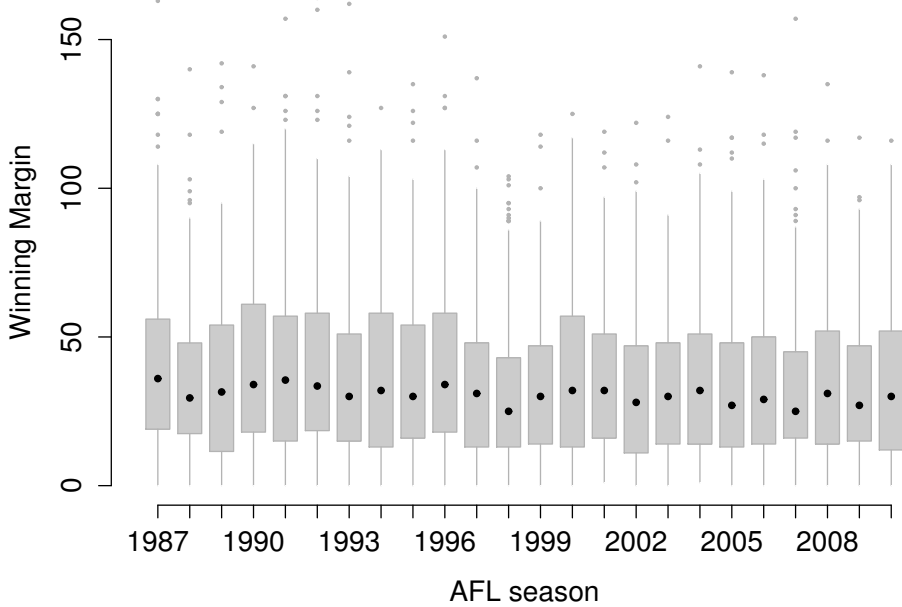


Figure 6.15: A cleaned up version of Figure 6.14. Notice that I've used a very minimalist design for the boxplots, so as to focus the eye on the medians. I've also converted the medians to solid dots, to convey a sense that year to year variation in the median should be thought of as a single coherent plot (similar to what we did when plotting the `Fibonacci` variable earlier). The size of outliers has been shrunk, because they aren't actually very interesting. In contrast, I've added a fill colour to the boxes, to make it easier to look at the trend in the interquartile range across years.

```

+      outpch = 20,          # outliers as solid dots
+      outcex = .5,          # shrink the outliers
+      medlty = "blank",     # no line for the medians
+      medpch = 20,          # instead, draw solid dots
+      medlwd = 1.5          # make them larger
+
)

```

Of course, given that the command is that long, you might have guessed that I didn't spend ages typing all that rubbish in over and over again. Instead, I wrote a script, which I kept tweaking until it produced the figure that I wanted. We'll talk about scripts later in Section 8.1, but given the length of the command I thought I'd remind you that there's an easier way of trying out different commands than typing them all in over and over.

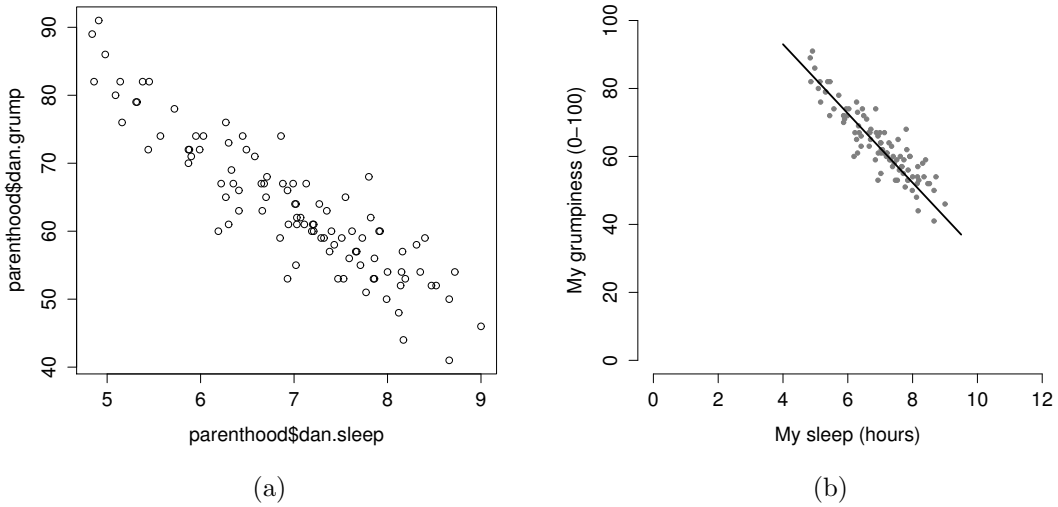


Figure 6.16: Two different scatterplots: (a) the default scatterplot that R produces, (b) one that makes use of several options for fancier display.

6.6

Scatterplots

Scatterplots are a simple but effective tool for visualising data. We've already seen scatterplots in this chapter, when using the `plot()` function to draw the `Fibonacci` variable as a collection of dots (Section 6.2). However, for the purposes of this section I have a slightly different notion in mind. Instead of just plotting one variable, what I want to do with my scatterplot is display the relationship between *two* variables, like we saw with the figures in the section on correlation (Section 5.7). It's this latter application that we usually have in mind when we use the term "scatterplot". In this kind of plot, each observation corresponds to one dot: the horizontal location of the dot plots the value of the observation on one variable, and the vertical location displays its value on the other variable. In many situations you don't really have a clear opinions about what the *causal* relationship is (e.g., does A cause B, or does B cause A, or does some other variable C control both A and B). If that's the case, it doesn't really matter which variable you plot on the x-axis and which one you plot on the y-axis. However, in many situations you do have a pretty strong idea which variable you think is most likely to be causal, or at least you have some suspicions in that direction. If so, then it's conventional to plot the cause variable on the x-axis, and the effect variable on the y-axis. With that in mind, let's look at how to draw scatterplots in R, using the same `parenthood` data set (i.e. `parenthood.Rdata`) that I used when introducing the idea of correlations.

Suppose my goal is to draw a scatterplot displaying the relationship between the amount of sleep that I get (`dan.sleep`) and how grumpy I am the next day (`dan.grump`). As you might expect given our earlier use of `plot()` to display the `Fibonacci` data, the function that we use is the `plot()` function, but because it's a generic function all the hard work is still being done by the `plot.default()` function. In any case, there are two different ways in which we can get the plot that we're after. The first way is to specify the name of the variable to be plotted on the `x` axis and the variable to be plotted on the `y` axis. When we

do it this way, the command looks like this:

```
> plot( x = parenthood$dan.sleep,    # data on the x-axis
+       y = parenthood$dan.grump     # data on the y-axis
+ )
```

The second way do to it is to use a “formula and data frame” format, but I’m going to avoid using it.¹³ For now, let’s just stick with the `x` and `y` version. If we do this, the result is the very basic scatterplot shown in Figure 6.16a. This serves fairly well, but there’s a few customisations that we probably want to make in order to have this work properly. As usual, we want to add some labels, but there’s a few other things we might want to do as well. Firstly, it’s sometimes useful to rescale the plots. In Figure 6.16a R has selected the scales so that the data fall neatly in the middle. But, in this case, we happen to know that the grumpiness measure falls on a scale from 0 to 100, and the hours slept falls on a natural scale between 0 hours and about 12 or so hours (the longest I can sleep in real life). So the command I might use to draw this is:

```
> plot( x = parenthood$dan.sleep,          # data on the x-axis
+       y = parenthood$dan.grump,          # data on the y-axis
+       xlab = "My sleep (hours)",        # x-axis label
+       ylab = "My grumpiness (0-100)",   # y-axis label
+       xlim = c(0,12),                  # scale the x-axis
+       ylim = c(0,100),                  # scale the y-axis
+       pch = 20,                       # change the plot type
+       col = "gray50",                 # dim the dots slightly
+       frame.plot = FALSE             # don't draw a box
+ )
```

This command produces the scatterplot in Figure 6.16b, or at least very nearly. What it doesn’t do is draw the line through the middle of the points. Sometimes it can be very useful to do this, and I can do so using `lines()`, which is a low level plotting function. Better yet, the arguments that I need to specify are pretty much the exact same ones that I use when calling the `plot()` function. That is, suppose that I want to draw a line that goes from the point (4,93) to the point (9.5,37). Then the `x` locations can be specified by the vector `c(4,9.5)` and the `y` locations correspond to the vector `c(93,37)`. In other words, I use this command:

```
> lines( x = c(4,9.5),    # the horizontal locations
+         y = c(93,37),    # the vertical locations
+         lwd = 2          # line width
+ )
```

And when I do so, R plots the line over the top of the plot that I drew using the previous command. In most realistic data analysis situations you absolutely don’t want to just guess where the line through the points goes, since there’s about a billion different ways in which you can get R to do a better job. However, it does at least illustrate the basic idea.

One possibility, if you do want to get R to draw nice clean lines through the data for you, is to use the `scatterplot()` function in the `car` package. Before we can use `scatterplot()` we need to load the package:

¹³The reason is that there’s an annoying design flaw in the way the `plot()` function handles this situation. The problem is that the `plot.formula()` function uses different names to for the arguments than the `plot()` function expects. As a consequence, you can’t specify the formula argument by name. If you just specify a formula as the first argument without using the name it works fine, because the `plot()` function thinks the formula corresponds to the `x` argument, and the `plot.formula()` function thinks it corresponds to the `formula` argument; and surprisingly, everything works nicely. But the moment that you, the user, tries to be unambiguous about the name, one of those two functions is going to cry.

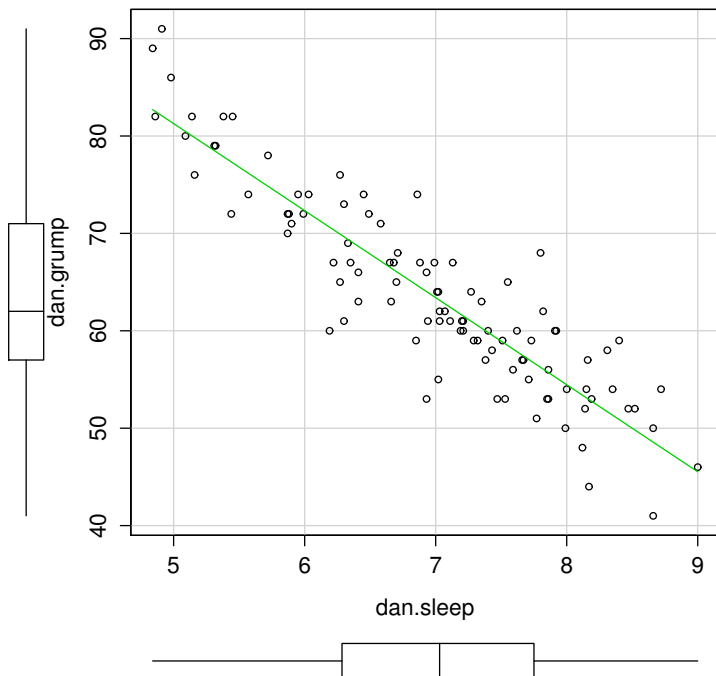


Figure 6.17: A fancy scatterplot drawn using the `scatterplot()` function in the `car` package.

```
> library( car )
```

Having done so, we can now use the function. The command we need is this one:

```
> scatterplot( dan.grump ~ dan.sleep,
+               data = parenthood,
+               smooth = FALSE
+ )
```

The first two arguments should be familiar: the first input is a formula (`dan.grump ~ dan.sleep`) telling R what variables to plot,¹⁴ and the second specifies a `data` frame. The third argument `smooth` I've set to `FALSE` to stop the `scatterplot()` function from drawing a fancy "smoothed" trendline (since it's a bit confusing to beginners). The scatterplot itself is shown in Figure 6.17. As you can see, it's not only drawn the scatterplot, but its also drawn boxplots for each of the two variables, as well as a simple line of best fit showing the relationship between the two variables.

¹⁴You might be wondering why I haven't specified the argument name for the formula. The reason is that there's a bug in how the `scatterplot()` function is written: under the hood there's one function that expects the argument to be named `x` and another one that expects it to be called `formula`. I don't know why the function was written this way, but it's not an isolated problem: this particular kind of bug repeats itself in a couple of other functions (you'll see it again in Chapter 13). The solution in such cases is to omit the argument name: that way, one function "thinks" that you've specified `x` and the other one "thinks" you've specified `formula` and everything works the way it's supposed to. It's not a great state of affairs, I'll admit, but it sort of works.

6.6.1 More elaborate options

Often you find yourself wanting to look at the relationships between several variables at once. One useful tool for doing so is to produce a **scatterplot matrix**, analogous to the correlation matrix.

```
> cor( x = parenthood ) # calculate correlation matrix
      dan.sleep  baby.sleep  dan.grump      day
dan.sleep    1.00000000  0.62794934 -0.90338404 -0.09840768
baby.sleep   0.62794934  1.00000000 -0.56596373 -0.01043394
dan.grump   -0.90338404 -0.56596373  1.00000000  0.07647926
day        -0.09840768 -0.01043394  0.07647926  1.00000000
```

We can get a the corresponding scatterplot matrix by using the **pairs()** function:¹⁵

```
> pairs( x = parenthood ) # draw corresponding scatterplot matrix
```

The output of the **pairs()** command is shown in Figure 6.18. An alternative way of calling the **pairs()** function, which can be useful in some situations, is to specify the variables to include using a one-sided formula. For instance, this

```
> pairs( formula = ~ dan.sleep + baby.sleep + dan.grump,
+         data = parenthood
+ )
```

would produce a 3×3 scatterplot matrix that only compare `dan.sleep`, `dan.grump` and `baby.sleep`. Obviously, the first version is much easier, but there are cases where you really only want to look at a few of the variables, so it's nice to use the formula interface.

6.7

Bar graphs

Another form of graph that you often want to plot is the **bar graph**. The main function that you can use in R to draw them is the **barplot()** function.¹⁶ And to illustrate the use of the function, I'll use the **finalists** variable that I introduced in Section 5.1.7. What I want to do is draw a bar graph that displays the number of finals that each team has played in over the time spanned by the **afl** data set. So, let's start by creating a vector that contains this information. I'll use the **tabulate()** function to do this (which will be discussed properly in Section 7.1), since it creates a simple numeric vector:

```
> freq <- tabulate( afl.finalists )
> print( freq )
[1] 26 25 26 28 32  0  6 39 27 28 28 17  6 24 26 39 24
```

This isn't exactly the prettiest of frequency tables, of course. I'm only doing it this way so that you can see the **barplot()** function in it's "purest" form: when the input is just an ordinary numeric vector. That being said, I'm obviously going to need the team names to create some labels, so let's create a variable

¹⁵ Yet again, we could have produced this output using the **plot()** function: when the **x** argument is a data frame containing numeric variables only, then the output is a scatterplot matrix. So, once again, what I could have done is just type **plot(parenthood)**.

¹⁶ Once again, it's worth noting the link to the generic **plot()** function. If the **x** argument to **plot()** is a factor (and no **y** argument is given), the result is a bar graph. So you could use **plot(afl.finalists)** and get the same output as **barplot(afl.finalists)**.

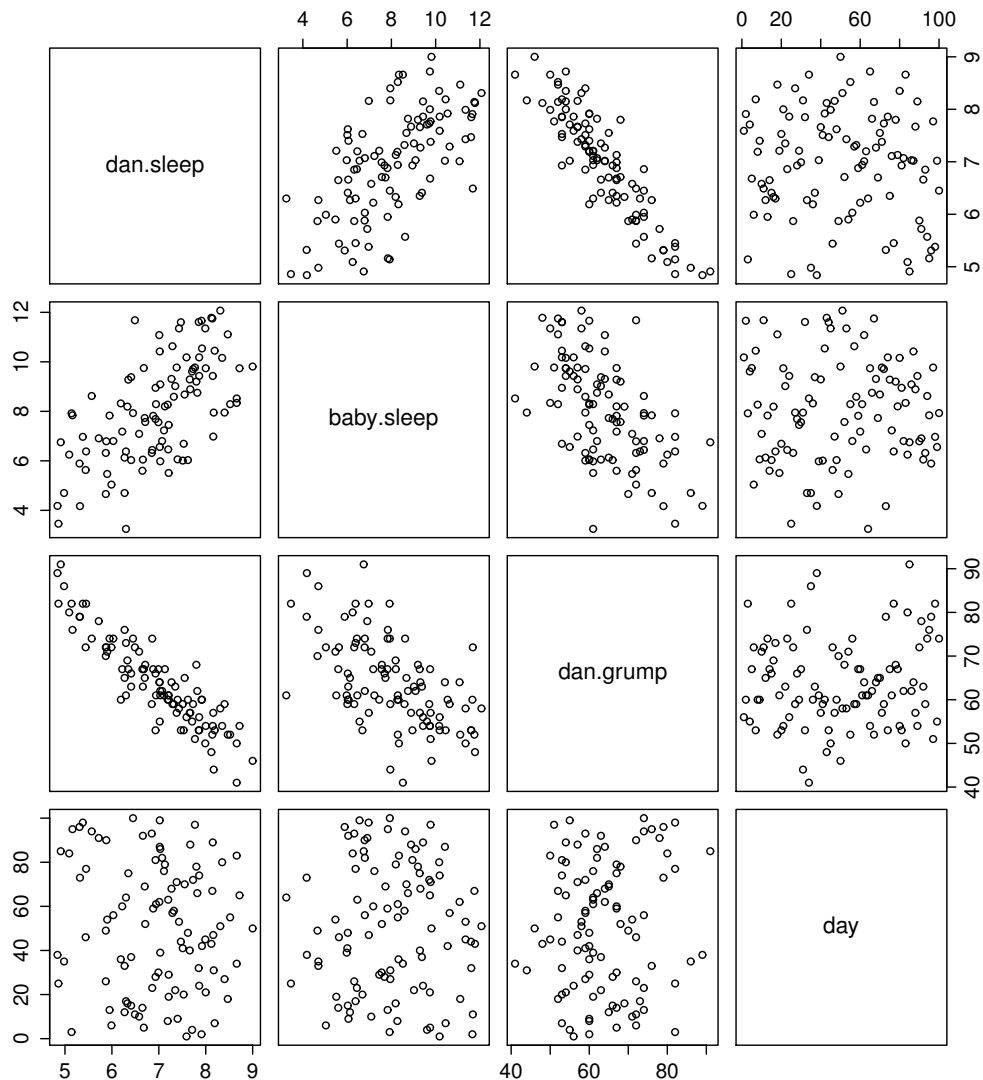


Figure 6.18: A matrix of scatterplots produced using `pairs()`.

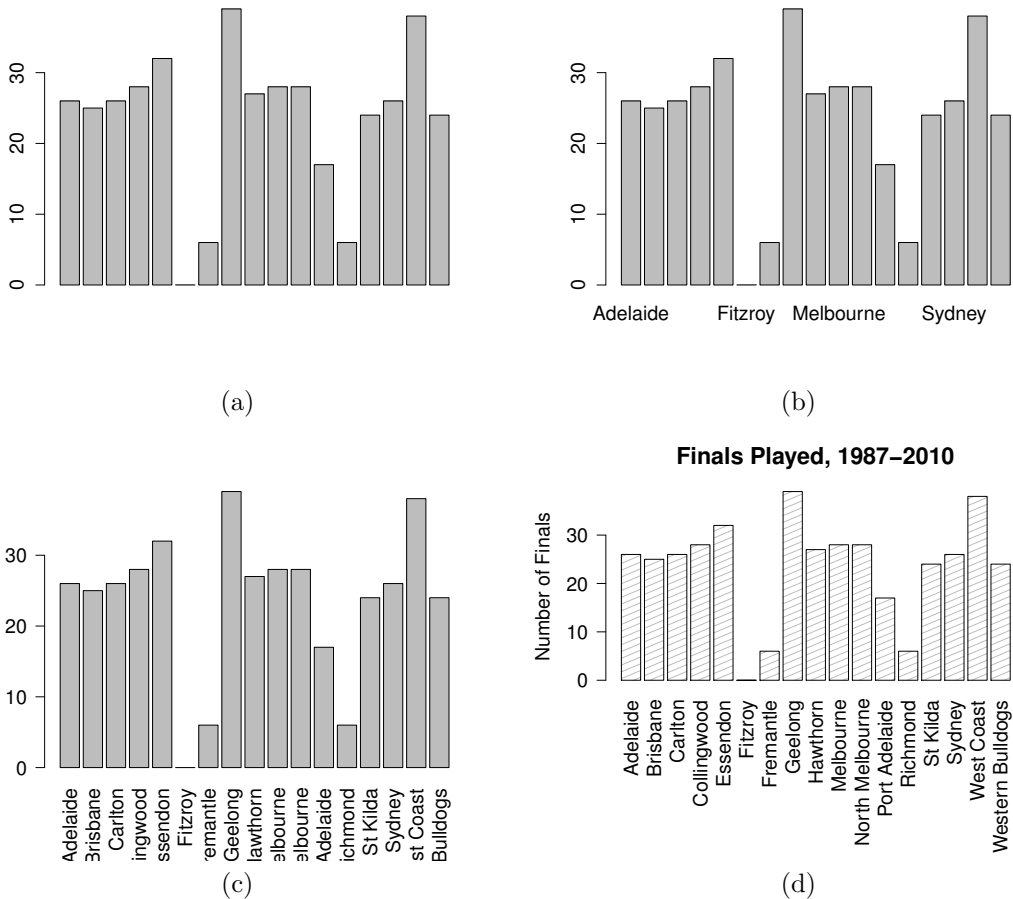


Figure 6.19: Four bargraphs. Panel a shows the simplest version of a bargraph, containing the data but no labels. In panel b we've added the labels, but because the text runs horizontally R only includes a few of them. In panel c we've rotated the labels, but now the text is too long to fit. Finally, in panel d we fix this by expanding the margin at the bottom, and add several other customisations to make the chart a bit nicer.

with those. I'll do this using the `levels()` function, which outputs the names of all the levels of a factor (see Section 4.7):

```
> teams <- levels( afl.finalists )
> print( teams )
[1] "Adelaide"          "Brisbane"           "Carlton"            "Collingwood"
[5] "Essendon"           "Fitzroy"             "Fremantle"          "Geelong"
[9] "Hawthorn"            "Melbourne"          "North Melbourne"    "Port Adelaide"
[13] "Richmond"           "St Kilda"            "Sydney"             "West Coast"
[17] "Western Bulldogs"
```

Okay, so now that we have the information we need, let's draw our bar graph. The main argument that you need to specify for a bar graph is the `height` of the bars, which in our case correspond to the values stored in the `freq` variable:

```
> barplot( height = freq ) # specifying the argument name (panel a)
> barplot( freq )        # the lazier version (panel a)
```

Either of these two commands will produce the simple bar graph shown in Figure 6.19a. As you can see, R has drawn a pretty minimal plot. It doesn't have any labels, obviously, because we didn't actually tell the `barplot()` function what the labels are! To do this, we need to specify the `names.arg` argument. The `names.arg` argument needs to be a vector of character strings containing the text that needs to be used as the label for each of the items. In this case, the `teams` vector is exactly what we need, so the command we're looking for is:

```
> barplot( height = freq, names.arg = teams ) # panel b
```

This is an improvement, but not much of an improvement. R has only included a few of the labels, because it can't fit them in the plot. This is the same behaviour we saw earlier with the multiple-boxplot graph in Figure 6.14. However, in Figure 6.14 it wasn't an issue: it's pretty obvious from inspection that the two unlabelled plots in between 1987 and 1990 must correspond to the data from 1988 and 1989. However, the fact that `barplot()` has omitted the names of every team in between Adelaide and Fitzroy is a lot more problematic.

The simplest way to fix this is to rotate the labels, so that the text runs vertically not horizontally. To do this, we need to alter set the `las` parameter, which I discussed briefly in Section 6.2. What I'll do is tell R to rotate the text so that it's always perpendicular to the axes (i.e., I'll set `las = 2`). When I do that, as per the following command...

```
> barplot( height = freq,      # the frequencies
+         names.arg = teams,   # the labels
+         las = 2              # rotate the labels
+ )                      # (see panel c)
```

... the result is the bar graph shown in Figure 6.19c. We've fixed the problem, but we've created a new one: the axis labels don't quite fit anymore. To fix this, we have to be a bit cleverer again. A simple fix would be to use shorter names rather than the full name of all teams, and in many situations that's probably the right thing to do. However, at other times you really do need to create a bit more space to add your labels, so I'll show you how to do that.

6.7.1 Changing global settings using `par()`

Altering the margins to the plot is actually a somewhat more complicated exercise than you might think. In principle it's a very simple thing to do: the size of the margins is governed by a graphical

parameter called `mar`, so all we need to do is alter this parameter. First, let's look at what the `mar` argument specifies. The `mar` argument is a vector containing four numbers: specifying the amount of space at the bottom, the left, the top and then the right. The units are “number of ‘lines’”. The default value for `mar` is `c(5.1, 4.1, 4.1, 2.1)`, meaning that R leaves 5.1 “lines” empty at the bottom, 4.1 lines on the left and the bottom, and only 2.1 lines on the right. In order to make more room at the bottom, what I need to do is change the first of these numbers. A value of 10.1 should do the trick.

So far this doesn't seem any different to the other graphical parameters that we've talked about. However, because of the way that the traditional graphics system in R works, you need to specify what the margins will be *before* calling your high-level plotting function. Unlike the other cases we've seen, you can't treat `mar` as if it were just another argument in your plotting function. Instead, you have to use the `par()` function to change the graphical parameters beforehand, and only then try to draw your figure. In other words, the first thing I would do is this:

```
> par( mar = c( 10.1, 4.1, 4.1, 2.1) )
```

There's no visible output here, but behind the scenes R has changed the graphical parameters associated with the current device (remember, in R terminology all graphics are drawn onto a “device”). Now that this is done, we could use the exact same command as before, but this time you'd see that the labels all fit, because R now leaves twice as much room for the labels at the bottom. However, since I've now figured out how to get the labels to display properly, I might as well play around with some of the other options, all of which are things you've seen before:

```
> barplot( height = freq,                                     # the data to plot
+           names.arg = teams,                                # label the plots
+           las = 2,                                         # rotate the labels
+           ylab = "Number of Finals",                         # y axis label
+           main = "Finals Played, 1987-2010",                # figure title
+           density = 10,                                      # shade the bars
+           angle = 20                                       # shading lines angle
+ )
```

However, one thing to remember about the `par()` function is that it doesn't just change the graphical parameters for the current *plot*. Rather, the changes pertain to any subsequent plot that you draw onto the same *device*. This might be exactly what you want, in which case there's no problem. But if not, you need to reset the graphical parameters to their original settings. To do this, you can either close the device (e.g., close the window, or click the “Clear All” button in the Plots pane in Rstudio) or you can reset the graphical parameters to their original values, using a command like this:

```
> par( mar = c(5.1, 4.1, 4.1, 2.1) )
```

6.8

Saving image files using R

Hold on, you might be thinking. What's the good of being able to draw pretty pictures in R if I can't save them and send them to friends to brag about how awesome my data is? How do I save the picture? To be honest, this is one of those situations where the easiest thing to do is to use the nice menu options on your system. Specifically:

- **Rstudio.** If you're running R through Rstudio, then the easiest way to save your image is to click on the “Export” button in the Plot panel (i.e., the area in Rstudio where all the plots have been

appearing). When you do that you'll see a menu that contains the options "Save Plot as PDF" and "Save Plot as Image". Either version works. Both will bring up dialog boxes that give you a few options that you can play with, but besides that it's pretty simple.

- **Windows.** If you're using the default application in Windows (i.e. Rgui.exe), then the way to save your image files is as follows. Assuming that you've got the graphics window on top, then you'll see four menus: "File", "History", "Resize" and "Windows". Just go to the "File" menu, then choose "Save as", and then choose what file type you'd like. It will then let you save the image.
- **Mac.** If you're using the default application in a Mac (i.e. R.app) the easiest way to get an image is also via the menus. Make sure the active window is the figure window (i.e., not the console), then click on the "File" menu, and choose "Save as". Unlike Windows, however, you don't get a lot of options for what file type you want: it's PDF or nothing! The easiest thing to do is to save it as a PDF. The reason I say this is that, on a Mac, it's very easy to convert PDFs to other image formats. All you have to do is open the PDF using the Mac "preview" app (i.e., don't use Acrobat reader). Once you've got it open in preview, go to the "File" menu and (again) choose "Save as". You can select among various file types.

This works pretty nicely for most situations. So, unless you're filled with a burning desire to learn the low level details, feel free to skip the rest of this section.

6.8.1 The ugly details

As I say, the menu-based options should be good enough for most people most of the time. However, one day you might want to be a bit more sophisticated, and make use of R's image writing capabilities at a lower level. In this section I'll give you a very basic introduction to this. In all honesty, this barely scratches the surface, but it will help a little bit in getting you started if you want to learn the details.

Okay, as I hinted earlier, whenever you're drawing pictures in R you're deemed to be drawing *to* a device of some kind. There are devices that correspond to a figure drawn on screen, and there are devices that correspond to graphics files that R will produce for you. For the purposes of this section I'll assume that you're using the default application in either Windows or Mac OS, not Rstudio. The reason for this is that my experience with the graphical device provided by Rstudio has led me to suspect that it still has a bunch on non-standard (or possibly just undocumented) features, and so I don't quite trust that it always does what I expect. I've no doubt they'll smooth it out later, but I can honestly say that I don't quite get what's going on with the `RStudioGD` device. In any case, we can ask R to list all of the graphics devices that currently exist, simply by using the command `dev.list()`. If there are no figure windows open, then you'll see this:

```
> dev.list()
NULL
```

which just means that R doesn't have any graphics devices open. However, suppose if you've just drawn a histogram and you type the same command, R will now give you a different answer. For instance, if you're using Windows:

```
> hist( afl.margins )
> dev.list()
windows
2
```

What this means is that there is one graphics device (device 2) that is currently open, and it's a figure window. If you did the same thing on a Mac, you get basically the same answer, except that the name

of the device would be `quartz` rather than `windows`. If you had several graphics windows open (which, incidentally, you can do by using the `dev.new()` command) then you'd see something like this:

```
> dev.list()
windows windows windows
      2      3      4
```

Okay, so that's the basic idea behind graphics devices. The key idea here is that graphics files (like JPEG images etc) are *also* graphics devices as far as R is concerned. So what you want to do is to *copy* the contents of one graphics device to another one. There's a command called `dev.copy()` that does this, but what I'll explain to you is a simpler one called `dev.print()`. It's pretty simple:

```
> dev.print( device = jpeg,           # what are we printing to?
+             filename = "thisfile.jpg", # name of the image file
+             width = 480,            # how many pixels wide should it be
+             height = 300            # how many pixels high should it be
+ )
```

This takes the “active” figure window, copies it to a jpeg file (which R treats as a device) and then closes that device. The `filename = "thisfile.jpg"` part tells R what to name the graphics file, and the `width = 480` and `height = 300` arguments tell R to draw an image that is 300 pixels high and 480 pixels wide. If you want a different kind of file, just change the device argument from `jpeg` to something else. R has devices for `png`, `tiff` and `bmp` that all work in exactly the same way as the `jpeg` command, but produce different kinds of files. Actually, for simple cartoonish graphics like this histogram, you'd be better advised to use PNG or TIFF over JPEG. The JPEG format is very good for natural images, but is wasteful for simple line drawings. The information above probably covers most things you might want to. However, if you want more information about what kinds of options you can specify using R, have a look at the help documentation by typing `?jpeg` or `?tiff` or whatever.

6.9

Summary

- Basic overview to R graphics (Section 6.1)
- An introduction to plotting (Section 6.2)
- Drawing histograms (Section 6.3)
- Producing stem and leaf plots (Section 6.4)
- Drawing boxplots (Section 6.5)
- Drawing scatterplots (Section 6.6)
- Drawing bar graphs (Section 6.7)
- Saving image files using R(Section 6.8)

7. Pragmatic matters

The garden of life never seems to confine itself to the plots philosophers have laid out for its convenience. Maybe a few more tractors would do the trick.

– Roger Zelazny

This is a somewhat strange chapter, even by my standards. My goal in this chapter is to talk a bit more honestly about the realities of working with data than you'll see anywhere else in the book. The problem with real world data sets is that they are *messy*. Very often the data file that you start out with doesn't have the variables stored in the right format for the analysis you want to do. Sometimes might be a lot of missing values in your data set. Sometimes you only want to analyse a subset of the data. Et cetera. In other words, there's a lot of **data manipulation** that you need to do, just to get all your data set into the format that you need it. The purpose of this chapter is to provide a basic introduction to all these pragmatic topics. Although the chapter is motivated by the kinds of practical issues that arise when manipulating real data, I'll stick with the practice that I've adopted through most of the book and rely on very small, toy data sets that illustrate the underlying issue. Because this chapter is essentially a collection of "tricks" and doesn't tell a single coherent story, it may be useful to start with a list of topics:

- Section 7.1. Tabulating data.
- Section 7.2. Transforming or recoding a variable.
- Section 7.3. Some useful mathematical functions.
- Section 7.4. Extracting a subset of a vector.
- Section 7.5. Extracting a subset of a data frame.
- Section 7.6. Sorting, flipping or merging data sets.
- Section 7.7. Reshaping a data frame.
- Section 7.8. Manipulating text.
- Section 7.9. Opening data from different file types.
- Section 7.10. Coercing data from one type to another.
- Section 7.11. Other important data types.
- Section 7.12. Miscellaneous topics.

As you can see, the list of topics that the chapter covers is pretty broad, and there's a *lot* of content there. Even though this is one of the longest and hardest chapters in the book, I'm really only scratching the surface of several fairly different and important topics. My advice, as usual, is to read through the chapter once and try to follow as much of it as you can. Don't worry too much if you can't grasp it all at once, especially the later sections. The rest of the book is only lightly reliant on this chapter, so you can

get away with just understanding the basics. However, what you'll probably find is that later on you'll need to flick back to this chapter in order to understand some of the concepts that I refer to here.

7.1

Tabulating and cross-tabulating data

A very common task when analysing data is the construction of frequency tables, or cross-tabulation of one variable against another. There are several functions that you can use in R for that purpose. In this section I'll illustrate the use of three functions – `table()`, `xtabs()` and `tabulate()` – though there are other options (e.g., `ftable()`) available.

7.1.1 Creating tables from vectors

Let's start with a simple example. As the father of a small child, I naturally spend a lot of time watching TV shows like *In the Night Garden*. In the `nightgarden.Rdata` file, I've transcribed a short section of the dialogue. The file contains two variables, `speaker` and `utterance`, and when we take a look at the data,

```
> load("nightgarden.Rdata")  
  
> who()  
-- Name -- -- Class -- -- Size --  
speaker      character      10  
utterance    character      10  
  
> print(speaker)  
[1] "upsy-daisy"  "upsy-daisy"  "upsy-daisy"  "upsy-daisy"  "tombliboo"  
[6] "tombliboo"   "makka-pakka" "makka-pakka" "makka-pakka" "makka-pakka"  
  
> print(utterance)  
[1] "pip" "pip" "onk" "onk" "ee"   "oo"   "pip" "pip" "onk" "onk"
```

it becomes very clear what happened to my sanity. With these as my data, one task I might find myself needing to do is construct a frequency count of the number of words each character speaks during the show. The `table()` function provides a simple way to do this. The basic usage of the `table()` function is as follows:

```
> table(speaker)  
speaker  
makka-pakka  tombliboo  upsy-daisy  
        4          2          4
```

The output here tells us on the first line that what we're looking at is a tabulation of the `speaker` variable. On the second line it lists all the different speakers that exist in the data, and on the third line it tells you how many times that speaker appears in the data. In other words, it's a frequency table¹. Notice that in the command above I didn't name the argument, since `table()` is another function that makes use of

¹As usual, you can assign this output to a variable. If you type `speaker.freq <- table(speaker)` at the command prompt R will store the table as a variable. If you then type `class(speaker.freq)` you'll see that the output is actually of class `table`. The key thing to note about a table object is that it's basically a matrix (see Section 7.11.1).

unnamed arguments. You just type in a list of the variables that you want R to tabulate, and it tabulates them. For instance, if I type in the name of two variables, what I get as the output is a cross-tabulation:

```
> table( speaker, utterance )
      utterance
speaker   ee onk oo pip
makka-pakka 0  2  0  2
tombliboo   1  0  1  0
upsy-daisy   0  2  0  2
```

When interpreting this table, remember that these are counts: so the fact that the first row and second column corresponds to a value of 2 indicates that Makka-Pakka (row 1) says “onk” (column 2) twice in this data set. As you’d expect, you can produce three way or higher order cross tabulations just by adding more objects to the list of inputs. However, I won’t discuss that in this section.

7.1.2 Creating tables from data frames

Most of the time your data are stored in a data frame, not kept as separate variables in the workspace. Let’s create one:

```
> itng <- data.frame( speaker, utterance )
> itng
  speaker utterance
1 upsy-daisy      pip
2 upsy-daisy      pip
3 upsy-daisy     onk
4 upsy-daisy     onk
5 tombliboo      ee
6 tombliboo      oo
7 makka-pakka    pip
8 makka-pakka    pip
9 makka-pakka    onk
10 makka-pakka   onk
```

There’s a couple of options under these circumstances. Firstly, if you just want to cross-tabulate all of the variables in the data frame, then it’s really easy:

```
> table( itng )
      utterance
speaker   ee onk oo pip
makka-pakka 0  2  0  2
tombliboo   1  0  1  0
upsy-daisy   0  2  0  2
```

However, it’s often the case that you want to select particular variables from the data frame to tabulate. This is where the `xtabs()` function is useful. In this function, you input a one sided `formula` in order to list all the variables you want to cross-tabulate, and the name of the `data` frame that stores the data:

```
> xtabs( formula = ~ speaker + utterance, data = itng )
      utterance
speaker   ee onk oo pip
makka-pakka 0  2  0  2
tombliboo   1  0  1  0
upsy-daisy   0  2  0  2
```

Clearly, this is a totally unnecessary command in the context of the `itng` data frame, but in most situations when you're analysing real data this is actually extremely useful, since your data set will almost certainly contain lots of variables and you'll only want to tabulate a few of them at a time.

7.1.3 Converting a table of counts to a table of proportions

The tabulation commands discussed so far all construct a table of raw frequencies: that is, a count of the total number of cases that satisfy certain conditions. However, often you want your data to be organised in terms of proportions rather than counts. This is where the `prop.table()` function comes in handy. It has two arguments:

- `x`. The frequency table that you want to convert.
- `margin`. Which “dimension” do you want to calculate proportions for. By default, R assumes you want the proportion to be expressed as a fraction of all possible events. See examples for details.

To see how this works:

```
> itng.table <- table( itng ) # create the table, and assign it to a variable
> itng.table # display the table again, as a reminder
      utterance
speaker    ee onk oo pip
makka-pakka 0   2   0   2
tombliboo   1   0   1   0
upsy-daisy  0   2   0   2

> prop.table( x = itng.table ) # express as proportion:
      utterance
speaker    ee onk oo pip
makka-pakka 0.0 0.2 0.0 0.2
tombliboo   0.1 0.0 0.1 0.0
upsy-daisy  0.0 0.2 0.0 0.2
```

Notice that there were 10 observations in our original data set, so all that R has done here is divide all our raw frequencies by 10. That's a sensible default, but more often you actually want to calculate the proportions separately by row (`margin = 1`) or by column (`margin = 2`). Again, this is most clearly seen by looking at examples:

```
> prop.table( x = itng.table, margin = 1 )
      utterance
speaker    ee onk oo pip
makka-pakka 0.0 0.5 0.0 0.5
tombliboo   0.5 0.0 0.5 0.0
upsy-daisy  0.0 0.5 0.0 0.5
```

Notice that each row now sums to 1, but that's not true for each column. What we're looking at here is the proportions of utterances made by each character. In other words, 50% of Makka-Pakka's utterances are “pip”, and the other 50% are “onk”. Let's contrast this with the following command:

```
> prop.table( x = itng.table, margin = 2 )
      utterance
speaker    ee onk oo pip
makka-pakka 0.0 0.5 0.0 0.5
tombliboo   1.0 0.0 1.0 0.0
upsy-daisy  0.0 0.5 0.0 0.5
```

Now the columns all sum to 1 but the rows don't. In this version, what we're seeing is the proportion of characters associated with each utterance. For instance, whenever the utterance "ee" is made (in this data set), 100% of the time it's a Tombliboo saying it.

7.1.4 Low level tabulation

One final function I want to mention is the `tabulate()` function, since this is actually the low-level function that does most of the hard work. It takes a numeric vector as input, and outputs frequencies as outputs:

```
> some.data <- c(1,2,3,1,1,3,1,1,2,8,3,1,2,4,2,3,5,2)
> tabulate(some.data)
[1] 6 5 4 1 1 0 0 1
```

7.2

Transforming and recoding a variable

It's not uncommon in real world data analysis to find that one of your variables isn't quite equivalent to the variable that you really want. For instance, it's often convenient to take a continuous-valued variable (e.g., age) and break it up into a smallish number of categories (e.g., younger, middle, older). At other times, you may need to convert a numeric variable into a different numeric variable (e.g., you may want to analyse at the absolute value of the original variable). In this section I'll describe a few key tricks that you can make use of to do this.

7.2.1 Creating a transformed variable

The first trick to discuss is the idea of **transforming** a variable. Taken literally, *anything* you do to a variable is a transformation, but in practice what it usually means is that you apply a relatively simple mathematical function to the original variable, in order to create new variable that either (a) provides a better way of describing the thing you're actually interested in or (b) is more closely in agreement with the assumptions of the statistical tests you want to do. Since – at this stage – I haven't talked about statistical tests or their assumptions, I'll show you an example based on the first case.

To keep the explanation simple, the variable we'll try to transform (`likert.raw`) isn't inside a data frame, though in real life it almost certainly would be. However, I think it's useful to start with an example that doesn't use data frames because it illustrates the fact that you already know how to do variable transformations. To see this, let's go through an example. Suppose I've run a short study in which I ask 10 people a single question:

On a scale of 1 (strongly disagree) to 7 (strongly agree), to what extent do you agree with the proposition that "Dinosaurs are awesome"?

Now let's load and look at the data. The data file `likert.Rdata` contains a single variable that contains the raw Likert-scale responses:

```
> load("likert.Rdata")
> likert.raw
[1] 1 7 3 4 4 4 2 6 5 5
```

However, if you think about it, this isn't the best way to represent these responses. Because of the fairly symmetric way that we set up the response scale, there's a sense in which the midpoint of the scale should have been coded as 0 (no opinion), and the two endpoints should be +3 (strong agree) and -3 (strong disagree). By recoding the data in this way, it's a bit more reflective of how we really think about the responses. The recoding here is trivially easy: we just subtract 4 from the raw scores:

```
> likert centred <- likert raw - 4
> likert centred
[1] -3 3 -1 0 0 0 -2 2 1 1
```

One reason why it might be useful to have the data in this format is that there are a lot of situations where you might prefer to analyse the *strength* of the opinion separately from the *direction* of the opinion. We can do two different transformations on this `likert centred` variable in order to distinguish between these two different concepts. Firstly, to compute an `opinion strength` variable, we want to take the absolute value of the centred data (using the `abs()` function that we've seen previously), like so:

```
> opinion strength <- abs( likert centred )
> opinion strength
[1] 3 3 1 0 0 0 2 2 1 1
```

Secondly, to compute a variable that contains only the direction of the opinion and ignores the strength, we can use the `sign()` function to do this. If you type `?sign` you'll see that this function is really simple: all negative numbers are converted to -1, all positive numbers are converted to 1 and zero stays as 0. So, when we apply the `sign()` function we obtain the following:

```
> opinion dir <- sign( likert centred )
> opinion dir
[1] -1 1 -1 0 0 0 -1 1 1 1
```

And we're done. We now have three shiny new variables, all of which are useful transformations of the original `likert raw` data. All of this should seem pretty familiar to you. The tools that you use to do regular calculations in R (e.g., Chapters 3 and 4) are very much the same ones that you use to transform your variables! To that end, in Section 7.3 I'll revisit the topic of doing calculations in R because there's a lot of other functions and operations that are worth knowing about.

Before moving on, you might be curious to see what these calculations look like if the data had started out in a data frame. To that end, it may help to note that the following example does all of the calculations using variables inside a data frame, and stores the variables created inside it:

```
> df <- data.frame( likert raw ) # create data frame
> df$likert centred <- df$likert raw - 4 # create centred data
> df$opinion strength <- abs( df$likert centred ) # create strength variable
> df$opinion dir <- sign( df$likert centred ) # create direction variable
> df # print the final data frame:

likert raw likert centred opinion strength opinion dir
1 1 -3 3 -1
2 7 3 3 1
3 3 -1 1 -1

BLAH BLAH BLAH
```

In other words, the commands you use are basically ones as before: it's just that every time you want to read a variable from the data frame or write to the data frame, you use the `$` operator. That's the easiest

way to do it, though I should make note of the fact that people sometimes make use of the `within()` function to do the same thing. However, since (a) I don't use the `within()` function anywhere else in this book, and (b) the `$` operator works just fine, I won't discuss it any further.

7.2.2 Cutting a numeric variable into categories

One pragmatic task that arises more often than you'd think is the problem of cutting a numeric variable up into discrete categories. For instance, suppose I'm interested in looking at the age distribution of people at a social gathering:

```
> age <- c( 60,58,24,26,34,42,31,30,33,2,9 )
```

In some situations it can be quite helpful to group these into a smallish number of categories. For example, we could group the data into three broad categories: young (0-20), adult (21-40) and older (41-60). This is a quite coarse-grained classification, and the labels that I've attached only make sense in the context of this data set (e.g., viewed more generally, a 42 year old wouldn't consider themselves as "older"). We can slice this variable up quite easily using the `cut()` function.² To make things a little cleaner, I'll start by creating a variable that defines the boundaries for the categories:

```
> age.breaks <- seq( from = 0, to = 60, by = 20 )
> age.breaks
[1] 0 20 40 60
```

and another one for the labels:

```
> age.labels <- c( "young", "adult", "older" )
> age.labels
[1] "young" "adult" "older"
```

Note that there are four numbers in the `age.breaks` variable, but only three labels in the `age.labels` variable; I've done this because the `cut()` function requires that you specify the *edges* of the categories rather than the mid-points. In any case, now that we've done this, we can use the `cut()` function to assign each observation to one of these three categories. There are several arguments to the `cut()` function, but the three that we need to care about are:

- `x`. The variable that needs to be categorised.
- `breaks`. This is either a vector containing the locations of the breaks separating the categories, or a number indicating how many categories you want.
- `labels`. The labels attached to the categories. This is optional: if you don't specify this R will attach a boring label showing the range associated with each category.

Since we've already created variables corresponding to the breaks and the labels, the command we need is just:

```
> age.group <- cut( x = age,                      # the variable to be categorised
+                     breaks = age.breaks,    # the edges of the categories
+                     labels = age.labels ) # the labels for the categories
```

²It's worth noting that there's also a more powerful function called `recode()` function in the `car` package that I won't discuss in this book but is worth looking into if you're looking for a bit more flexibility.

Note that the output variable here is a factor. In order to see what this command has actually done, we could just print out the `age.group` variable, but I think it's actually more helpful to create a data frame that includes both the original variable and the categorised one, so that you can see the two side by side:

```
> data.frame( age, age.group )
   age age.group
1   60      older
2   58      older
3   24     adult

BLAH BLAH BLAH

10    2      young
11    9      young
```

It can also be useful to tabulate the output, just to see if you've got a nice even division of the sample:

```
> table( age.group )
age.group
young adult older
2       6       3
```

In the example above, I made all the decisions myself. Much like the `hist()` function that we saw in Chapter 6, if you want to you can delegate a lot of the choices to R. For instance, if you want you can specify the *number* of categories you want, rather than giving explicit ranges for them, and you can allow R to come up with some labels for the categories. To give you a sense of how this works, have a look at the following example:

```
> age.group2 <- cut( x = age, breaks = 3 )
```

With this command, I've asked for three categories, but let R make the choices for where the boundaries should be. I won't bother to print out the `age.group2` variable, because it's not terribly pretty or very interesting. Instead, all of the important information can be extracted by looking at the tabulated data:

```
> table( age.group2 )
age.group2
(1.94,21.3] (21.3,40.7] (40.7,60.1]
2             6             3
```

This output takes a little bit of interpretation, but it's not complicated. What R has done is determined that the lowest age category should run from 1.94 years up to 21.3 years, the second category should run from 21.3 years to 40.7 years, and so on. The formatting on those labels might look a bit funny to those of you who haven't studied a lot of maths, but it's pretty simple. When R describes the first category as corresponding to the range (1.94, 21.3] what it's saying is that the range consists of those numbers that are larger than 1.94 but less than *or equal to* 21.3. In other words, the weird asymmetric brackets is R's way of telling you that if there happens to be a value that is exactly equal to 21.3, then it belongs to the first category, not the second one. Obviously, this isn't actually possible since I've only specified the ages to the nearest whole number, but R doesn't know this and so it's trying to be precise just in case. This notation is actually pretty standard, but I suspect not everyone reading the book will have seen it before. In any case, those labels are pretty ugly, so it's usually a good idea to specify your own, meaningful labels to the categories.

Before moving on, I should take a moment to talk a little about the mechanics of the `cut()` function. Notice that R has tried to divide the `age` variable into three roughly equal sized bins. Unless you specify

the particular breaks you want, that's what it will do. But suppose you want to divide the `age` variable into three categories of different size, but with approximately identical numbers of people. How would you do that? Well, if that's the case, then what you want to do is have the breaks correspond to the 0th, 33rd, 66th and 100th percentiles of the data. One way to do this would be to calculate those values using the `quantiles()` function and then use those quantiles as input to the `cut()` function. That's pretty easy to do, but it does take a couple of lines to type. So instead, the `lsr` package has a function called `quantileCut()` that does exactly this:

```
> age.group3 <- quantileCut( x = age, n = 3 )
> table( age.group3 )
age.group3
(1.94,27.3] (27.3,33.7] (33.7,60.1]
        4            3            4
```

Notice the difference in the boundaries that the `quantileCut()` function selects. The first and third categories now span an age range of about 25 years each, whereas the middle category has shrunk to a span of only 6 years. There are some situations where this is genuinely what you want (that's why I wrote the function!), but in general you should be careful. Usually the numeric variable that you're trying to cut into categories is already expressed in meaningful units (i.e., it's interval scale), but if you cut it into unequal bin sizes then it's often very difficult to attach meaningful interpretations to the resulting categories.

More generally, regardless of whether you're using the original `cut()` function or the `quantileCut()` version, it's important to take the time to figure out whether or not the resulting categories make any sense at all in terms of your research project. If they don't make any sense to you as meaningful categories, then any data analysis that uses those categories is likely to be just as meaningless. More generally, in practice I've noticed that people have a very strong desire to carve their (continuous and messy) data into a few (discrete and simple) categories; and then run analysis using the categorised data instead of the original one.³ I wouldn't go so far as to say that this is an inherently bad idea, but it does have some fairly serious drawbacks at times, so I would advise some caution if you are thinking about doing it.

7.3

A few more mathematical functions and operations

In Section 7.2 I discussed the ideas behind variable transformations, and showed that a lot of the transformations that you might want to apply to your data are based on fairly simple mathematical functions and operations, of the kind that we discussed in Chapter 3. In this section I want to return to that discussion, and mention several other mathematical functions and arithmetic operations that I didn't bother to mention when introducing you to R, but are actually quite useful for a lot of real world data analysis. Table 7.1 gives a brief overview of the various mathematical functions I want to talk about (and some that I already have talked about). Obviously this doesn't even come close to cataloguing the range of possibilities available in R, but it does cover a very wide range of functions that are used in day to day data analysis.

³If you've read further into the book, and are re-reading this section, then a good example of this would be someone choosing to do an ANOVA using `age.group3` as the grouping variable, instead of running a regression using `age` as a predictor. There are sometimes good reasons for doing this: for instance, if the relationship between `age` and your outcome variable is highly non-linear, and you aren't comfortable with trying to run non-linear regression! However, unless you really do have a good rationale for doing this, it's best not to. It tends to introduce all sorts of other problems (e.g., the data will probably violate the normality assumption), and you can lose a lot of power.

Table 7.1: Some of the mathematical functions available in R.

	function	example input	(answer)
square root	<code>sqrt()</code>	<code>sqrt(25)</code>	5
absolute value	<code>abs()</code>	<code>abs(-23)</code>	23
logarithm (base 10)	<code>log10()</code>	<code>log10(1000)</code>	3
logarithm (base e)	<code>log()</code>	<code>log(1000)</code>	6.908
exponentiation	<code>exp()</code>	<code>exp(6.908)</code>	1000.245
rounding to nearest	<code>round()</code>	<code>round(1.32)</code>	1
rounding down	<code>floor()</code>	<code>floor(1.32)</code>	1
rounding up	<code>ceiling()</code>	<code>ceiling(1.32)</code>	2

7.3.1 Rounding a number

One very simple transformation that crops up surprisingly often is the need to round a number to the nearest whole number, or to a certain number of significant digits. To start with, let's assume that we want to round to a whole number. To that end, there are three useful functions in R you want to know about: `round()`, `floor()` and `ceiling()`. The `round()` function just rounds to the *nearest* whole number. So if you round the number 4.3, it “rounds down” to 4, like so:

```
> round( x = 4.3 )
[1] 4
```

In contrast, if we want to round the number 4.7, we would round upwards to 5. In everyday life, when someone talks about “rounding”, they usually mean “round to nearest”, so this is the function we use most of the time. However sometimes you have reasons to want to always round up or always round down. If you want to always round down, use the `floor()` function instead; and if you want to force R to round up, then use `ceiling()`. That's the only difference between the three functions. What if you want to round to a certain number of digits? Let's suppose you want to round to a fixed number of decimal places, say 2 decimal places. If so, what you need to do is specify the `digits` argument to the `round()` function. That's pretty straightforward:

```
> round( x = 0.0123, digits = 2 )
[1] 0.01
```

The only subtlety that you need to keep in mind is that sometimes what you want to do is round to 2 **significant digits** and not to two decimal places. The difference is that, when determining the number of significant digits, zeros don't count. To see this, let's apply the `signif()` function instead of the `round()` function:

```
> signif( x = 0.0123, digits = 2 )
[1] 0.012
```

This time around, we get an answer of 0.012 because the zeros don't count as significant digits. Quite often scientific journals will ask you to report numbers to two or three significant digits, so it's useful to remember the distinction.

Table 7.2: Two more arithmetic operations that sometimes come in handy.

operation	operator	example input	example output
integer division	<code>%/%</code>	<code>42 %/% 10</code>	4
modulus	<code>%%</code>	<code>42 %% 10</code>	2

7.3.2 Modulus and integer division

Since we're on the topic of simple calculations, there are two other arithmetic operations that I should mention, since they can come in handy when working with real data. These operations are calculating a modulus and doing integer division. They don't come up anywhere else in this book, but they are worth knowing about. First, let's consider **integer division**. Suppose I have \$42 in my wallet, and want to buy some sandwiches, which are selling for \$10 each. How many sandwiches can I afford⁴ to buy? The answer is of course 4. Note that it's not 4.2, since no shop will sell me one-fifth of a sandwich. That's integer division. In R we perform integer division by using the `%/%` operator:

```
> 42 %/% 10
[1] 4
```

Okay, that's easy enough. What about the **modulus**? Basically, a modulus is the remainder after integer division, and it's calculated using the `%%` operator. For the sake of argument, let's suppose I buy four overpriced \$10 sandwiches. If I started out with \$42, how much money do I have left? The answer, as both R and common sense tells us, is \$2:

```
> 42 %% 10
[1] 2
```

So that's also pretty easy. There is, however, one subtlety that I need to mention, and this relates to how negative numbers are handled. Firstly, what would happen if I tried to do integer division with a negative number? Let's have a look:

```
> -42 %/% 10
[1] -5
```

This might strike you as counterintuitive: why does `42 %/% 10` produce an answer of 4, but `-42 %/% 10` gives us an answer of -5? Intuitively you might think that the answer to the second one should be -4. The way to think about it is like this. Suppose I *owe* the sandwich shop \$42, but I don't have any money. How many sandwiches would *I* have to give *them* in order to stop them from calling security? The answer⁵ here is 5, not 4. If I handed them 4 sandwiches, I'd still owe them \$2, right? So I actually have to give them 5 sandwiches. And since it's *me* giving them the sandwiches, the answer to `-42 %/% 10` is -5. As you might expect, the behaviour of the modulus operator has a similar pattern. If I've handed 5 sandwiches over to the shop in order to pay off my debt of \$42, then *they* now owe me \$8. So the modulus is now:

```
> -42 %% 10
[1] 8
```

⁴The real answer is 0: \$10 for a sandwich is a total ripoff so I should go next door and buy noodles.

⁵Again, I doubt that's the right "real world" answer. I suspect that most sandwich shops won't allow you to pay off your debts to them in sandwiches. But you get the idea.

7.3.3 Logarithms and exponentials

As I've mentioned earlier, R has an incredible range of mathematical functions built into it, and there really wouldn't be much point in trying to describe or even list all of them. For the most part, I've focused only on those functions that are strictly necessary for this book. However I do want to make an exception for logarithms and exponentials. Although they aren't needed anywhere else in this book, they are *everywhere* in statistics more broadly, and not only that, there are a *lot* of situations in which it is convenient to analyse the logarithm of a variable (i.e., to take a "log-transform" of the variable). I suspect that many (maybe most) readers of this book will have encountered logarithms and exponentials before, but from past experience I know that there's a substantial proportion of students who take a social science statistics class who haven't touched logarithms since high school, and would appreciate a bit of a refresher.

In order to understand logarithms and exponentials, the easiest thing to do is to actually calculate them and see how they relate to other simple calculations. There are three R functions in particular that I want to talk about, namely `log()`, `log10()` and `exp()`. To start with, let's consider `log10()`, which is known as the "logarithm in base 10". The trick to understanding a **logarithm** is to understand that it's basically the "opposite" of taking a power. Specifically, the logarithm in base 10 is closely related to the powers of 10. So let's start by noting that $10^3 = 1000$. Mathematically, we would write this:

$$10^3 = 1000$$

and in R we'd calculate it by using the command `10^3`. The trick to understanding a logarithm is to recognise that the statement that "10 to the power of 3 is equal to 1000" is equivalent to the statement that "the logarithm (in base 10) of 1000 is equal to 3". Mathematically, we write this as follows,

$$\log_{10}(1000) = 3$$

and if we wanted to do the calculation in R we would type this:

```
> log10( 1000 )
[1] 3
```

Obviously, since you already know that $10^3 = 1000$ there's really no point in getting R to tell you that the base-10 logarithm of 1000 is 3. However, most of the time you probably don't know what right answer is. For instance, I can honestly say that I didn't know that $10^{2.69897} = 500$, so it's rather convenient for me that I can use R to calculate the base-10 logarithm of 500:

```
> log10( 500 )
[1] 2.69897
```

Or at least it would be convenient if I had a pressing need to know the base-10 logarithm of 500.

Okay, since the `log10()` function is related to the powers of 10, you might expect that there are other logarithms (in bases other than 10) that are related to other powers too. And of course that's true: there's not really anything mathematically special about the number 10. You and I happen to find it useful because decimal numbers are built around the number 10, but the big bad world of mathematics scoffs at our decimal numbers. Sadly, the universe doesn't actually care how we write down numbers. Anyway, the consequence of this cosmic indifference is that there's nothing particularly special about calculating logarithms in base 10. You could, for instance, calculate your logarithms in base 2, and in fact R does provide a function for doing that, which is (not surprisingly) called `log2()`. Since we know that $2^3 = 2 \times 2 \times 2 = 8$, it's not surprise to see that

```
> log2( 8 )
[1] 3
```

Alternatively, a third type of logarithm – and one we see a lot more of in statistics than either base 10 or base 2 – is called the **natural logarithm**, and corresponds to the logarithm in base e . Since you might one day run into it, I'd better explain what e is. The number e , known as **Euler's number**, is one of those annoying “irrational” numbers whose decimal expansion is infinitely long, and is considered one of the most important numbers in mathematics. The first few digits of e are:

$$e = 2.718282$$

There are quite a few situations in statistics that require us to calculate powers of e , though none of them appear in this book. Raising e to the power x is called the **exponential** of x , and so it's very common to see e^x written as `exp(x)`. And so it's no surprise that R has a function that calculates exponentials, called `exp()`. For instance, suppose I wanted to calculate e^3 . I could try typing in the value of e manually, like this:

```
> 2.718282 ^ 3  
[1] 20.08554
```

but it's much easier to do the same thing using the `exp()` function:

```
> exp( 3 )  
[1] 20.08554
```

Anyway, because the number e crops up so often in statistics, the natural logarithm (i.e., logarithm in base e) also tends to turn up. Mathematicians often write it as $\log_e(x)$ or $\ln(x)$, or sometimes even just $\log(x)$. In fact, R works the same way: the `log()` function corresponds to the natural logarithm⁶. Anyway, as a quick check, let's calculate the natural logarithm of 20.08554 using R:

```
> log( 20.08554 )  
[1] 3
```

And with that, I think we've had quite enough exponentials and logarithms for this book!

7.4

Extracting a subset of a vector

One very important kind of data handling is being able to extract a particular subset of the data. For instance, you might be interested only in analysing the data from one experimental condition, or you may want to look closely at the data from people over 50 years in age. To do this, the first step is getting R to extract the subset of the data corresponding to the observations that you're interested in. In this section I'll talk about subsetting as it applies to vectors, extending the discussion from Chapters 3 and 4. In Section 7.5 I'll go on to talk about how this discussion extends to data frames.

7.4.1 Refresher

This section returns to the `nightgarden.Rdata` data set. If you're reading this whole chapter in one sitting, then you should already have this data set loaded. If not, don't forget to use the `load("nightgarden.Rdata")`

⁶Actually, that's a bit of a lie: the `log()` function is more flexible than that, and can be used to calculate logarithms in *any* base. The `log()` function has a `base` argument that you can specify, which has a default value of e . Thus `log10(1000)` is actually equivalent to `log(x = 1000, base = 10)`.

command. For this section, let's ignore the `itng` data frame that we created earlier, and focus instead on the two vectors `speaker` and `utterance` (see Section 7.1 if you've forgotten what those vectors look like). Suppose that what I want to do is pull out only those utterances that were made by Makka-Pakka. To that end, I could first use the equality operator to have R tell me which cases correspond to Makka-Pakka speaking:

```
> is.MP.speaking <- speaker == "makka-pakka"
> is.MP.speaking
[1] FALSE FALSE FALSE FALSE FALSE FALSE TRUE TRUE TRUE TRUE
```

and then use logical indexing to get R to print out those elements of `utterance` for which `is.MP.speaking` is true, like so:

```
> utterance[ is.MP.speaking ]
[1] "pip" "pip" "onk" "onk"
```

Or, since I'm lazy, I could collapse it to a single command like so:

```
> utterance[ speaker == "makka-pakka" ]
[1] "pip" "pip" "onk" "onk"
```

7.4.2 Using `%in%` to match multiple cases

A second useful trick to be aware of is the `%in%` operator⁷. It's actually very similar to the `==` operator, except that you can supply a collection of acceptable values. For instance, suppose I wanted to keep only those cases when the utterance is either "pip" or "oo". One simple way do to this is:

```
> utterance %in% c("pip", "oo")
[1] TRUE TRUE FALSE FALSE FALSE TRUE TRUE TRUE FALSE FALSE
```

What this does if return `TRUE` for those elements of `utterance` that are either "pip" or "oo" and returns `FALSE` for all the others. What that means is that if I want a list of all those instances of characters speaking either of these two words, I could do this:

```
> speaker[ utterance %in% c("pip", "oo") ]
[1] "upsy-daisy" "upsy-daisy" "tombliboo" "makka-pakka" "makka-pakka"
```

7.4.3 Using negative indices to drop elements

Before moving onto data frames, there's a couple of other tricks worth mentioning. The first of these is to use negative values as indices. Recall from Section 3.9 that we can use a vector of numbers to extract a set of elements that we would like to keep. For instance, suppose I want to keep only elements 2 and 3 from `utterance`. I could do so like this:

```
> utterance[2:3]
[1] "pip" "onk"
```

But suppose, on the other hand, that I have discovered that observations 2 and 3 are untrustworthy, and I want to keep everything *except* those two elements. To that end, R lets you use negative numbers to remove specific values, like so:

⁷It's also worth checking out the `match()` function

```
> utterance [ -(2:3) ]
[1] "pip" "onk" "ee" "oo" "pip" "pip" "onk" "onk"
```

The output here corresponds to element 1 of the original vector, followed by elements 4, 5, and so on. When all you want to do is remove a few cases, this is a very handy convention.

7.4.4 Splitting a vector by group

One particular example of subsetting that is especially common is the problem of splitting one variable up into several different variables, one corresponding to each group. For instance, in our *In the Night Garden* example, I might want to create subsets of the `utterance` variable for every character. One way to do this would be to just repeat the exercise that I went through earlier separately for each character, but that quickly gets annoying. A faster way to do it is to use the `split()` function. The arguments are:

- `x`. The variable that needs to be split into groups.
- `f`. The grouping variable.

What this function does is output a list (Section 4.9), containing one variable for each group. For instance, I could split up the `utterance` variable by `speaker` using the following command:

```
> speech.by.char <- split( x = utterance, f = speaker )
> speech.by.char
$`makka-pakka`
[1] "pip" "pip" "onk" "onk"

$tombliboo
[1] "ee" "oo"

$`upsy-daisy`
[1] "pip" "pip" "onk" "onk"
```

Once you're starting to become comfortable working with lists and data frames, this output is all you need, since you can work with this list in much the same way that you would work with a data frame. For instance, if you want the first utterance made by Makka-Pakka, all you need to do is type this:

```
> speech.by.char$`makka-pakka`[1]
[1] "pip"
```

Just remember that R does need you to add the quoting characters (i.e. `'`). Otherwise, there's nothing particularly new or difficult here.

However, sometimes – especially when you're just starting out – it can be convenient to pull these variables out of the list, and into the workspace. This isn't too difficult to do, though it can be a little daunting to novices. To that end, I've included a function called `importList()` in the `lsr` package that does this.⁸ First, here's what you'd have if you had wiped the workspace before the start of this section:

```
> who()
-- Name --      -- Class --      -- Size --
```

⁸It also works on data frames if you ever feel the need to import all of your variables from the data frame into the workspace. This can be useful at times, though it's not a good idea if you have large data sets or if you're working with multiple data sets at once. In particular, if you do this, never forget that you now have *two* copies of all your variables, one in the workspace and another in the data frame.

```

speaker      character   10
speech.by.char  list       3
utterance    character   10

```

Now we use the `importList()` function to copy all of the variables within the `speech.by.char` list:

```

> importList( speech.by.char )
Names of variables to be created:
[1] "makka.pakka" "tombliboo"   "upsy.daisy"
Create these variables? [y/n]

```

Because the `importList()` function is attempting to create new variables based on the names of the elements of the list, it pauses to check that you're okay with the variable names. The reason it does this is that, if one of the to-be-created variables has the same name as a variable that you already have in your workspace, that variable will end up being overwritten, so it's a good idea to check. Assuming that you type `y`, it will go on to create the variables. Nothing *appears* to have happened, but if we look at our workspace now:

```

> who()
-- Name --      -- Class --      -- Size --
makka.pakka    character     4
speaker         character    10
speech.by.char  list        3
tombliboo       character     2
upsy.daisy      character     4
utterance        character    10

```

we see that there are three new variables, called `makka.pakka`, `tombliboo` and `upsy.daisy`. Notice that the `importList()` function has converted the original character strings into valid R variable names, so the variable corresponding to `"makka-pakka"` is actually `makka.pakka`.⁹ Nevertheless, even though the names can change, note that each of these variables contains the exact same information as the original elements of the list did. For example:

```

> makka.pakka
[1] "pip" "pip" "onk" "onk"

```

7.5

Extracting a subset of a data frame

In this section we turn to the question of how to subset a data frame rather than a vector. To that end, the first thing I should point out is that, if all you want to do is subset *one* of the variables inside the data frame, then as usual the `$` operator is your friend. For instance, suppose I'm working with the `itng` data frame, and what I want to do is create the `speech.by.char` list. I can use the exact same tricks that I used last time, since what I really want to do is `split()` the `itng$utterance` vector, using the `itng$speaker` vector as the grouping variable. However, most of the time what you actually want to do is select several different variables within the data frame (i.e., keep only some of the columns), or maybe a subset of cases (i.e., keep only some of the rows). In order to understand how this works, we need to talk more specifically about data frames and how to subset them.

⁹You can do this yourself using the `make.names()` function. In fact, this is itself a handy thing to know about. For example, if you want to convert the names of the variables in the `speech.by.char` list into valid R variable names, you could use a command like this: `names(speech.by.char) <- make.names(names(speech.by.char))`. However, I won't go into details here.

7.5.1 Using the `subset()` function

There are several different ways to subset a data frame in R, some easier than others. I'll start by discussing the `subset()` function, which is probably the conceptually simplest way to do it. For our purposes there are three different arguments that you'll be most interested in:

- `x`. The data frame that you want to subset.
- `subset`. A vector of logical values indicating which cases (rows) of the data frame you want to keep. By default, all cases will be retained.
- `select`. This argument indicates which variables (columns) in the data frame you want to keep. This can either be a list of variable names, or a logical vector indicating which ones to keep, or even just a numeric vector containing the relevant column numbers. By default, all variables will be retained.

Let's start with an example in which I use all three of these arguments. Suppose that I want to subset the `itng` data frame, keeping only the utterances made by Makka-Pakka. What that means is that I need to use the `select` argument to pick out the `utterance` variable, and I also need to use the `subset` variable, to pick out the cases when Makka-Pakka is speaking (i.e., `speaker == "makka-pakka"`). Therefore, the command I need to use is this:

```
> df <- subset( x = itng,                                     # data frame is itng
+                 subset = speaker == "makka-pakka",      # keep only Makka-Pakkas speech
+                 select = utterance )                   # keep only the utterance variable

> print( df )
  utterance
7     pip
8     pip
9     onk
10    onk
```

The variable `df` here is still a data frame, but it only contains one variable (called `utterance`) and four cases. Notice that the row numbers are actually the same ones from the original data frame. It's worth taking a moment to briefly explain this. The reason that this happens is that these "row numbers" are actually row *names*. When you create a new data frame from scratch R will assign each row a fairly boring row name, which is identical to the row number. However, when you subset the data frame, each row keeps its original row name. This can be quite useful, since – as in the current example – it provides you a visual reminder of what each row in the new data frame corresponds to in the original data frame. However, if it annoys you, you can change the row names using the `rownames()` function.¹⁰

In any case, let's return to the `subset()` function, and look at what happens when we don't use all three of the arguments. Firstly, suppose that I didn't bother to specify the `select` argument. Let's see what happens:

```
> subset( x = itng,
+           subset = speaker == "makka-pakka" )

  speaker utterance
7 makka-pakka     pip
8 makka-pakka     pip
9 makka-pakka     onk
10 makka-pakka    onk
```

¹⁰Conveniently, if you type `rownames(df) <- NULL` R will renumber all the rows from scratch. For the `df` data frame, the labels that currently run from 7 to 10 will be changed to go from 1 to 4.

Not surprisingly, R has kept the same cases from the original data set (i.e., rows 7 through 10), but this time it has kept all of the variables from the data frame. Equally unsurprisingly, if I don't specify the `subset` argument, what we find is that R keeps all of the cases:

```
> subset( x = itng,
+         select = utterance )

      utterance
1       pip
2       pip
3       onk

BLAH BLAH BLAH
```

where the `BLAH BLAH BLAH` in this case shows all 10 cases from the original `itng` data frame. Again, it's important to note that this output is still a data frame: it's just a data frame with only a single variable.

7.5.2 Using square brackets: I. Rows and columns

Throughout the book so far, whenever I've been subsetting a vector I've tended use the square brackets `[]` to do so. But in the previous section when I started talking about subsetting a data frame I used the `subset()` function. As a consequence, you might be wondering whether it is possible to use the square brackets to subset a data frame. The answer, of course, is yes. Not only *can* you use square brackets for this purpose, as you become more familiar with R you'll find that this is actually much more convenient than using `subset()`. Unfortunately, the use of square brackets for this purpose is somewhat complicated, and can be very confusing to novices. So be warned: this section is more complicated than it feels like it "should" be. With that warning in place, I'll try to walk you through it slowly. For this section, I'll use a slightly different data set, namely the `garden` data frame that is stored in the "`nightgarden2.Rdata`" file.

```
> load( "nightgarden2.Rdata" )
> garden
      speaker utterance line
case.1  upsy-daisy     pip    1
case.2  upsy-daisy     pip    2
case.3  tombliboo      ee     5
case.4  makka-pakka   pip    7
case.5  makka-pakka   onk    9
```

As you can see, the `garden` data frame contains 3 variables and 5 cases, and this time around I've used the `rownames()` function to attach slightly verbose labels to each of the cases. Moreover, let's assume that what we want to do is to pick out rows 4 and 5 (the two cases when Makka-Pakka is speaking), and columns 1 and 2 (variables `speaker` and `utterance`).

How shall we do this? As usual, there's more than one way. The first way is based on the observation that, since a data frame is basically a table, every element in the data frame has a row number and a column number. So, if we want to pick out a single element, we have to specify the row number *and* a column number within the square brackets. By convention, the row number comes first. So, for the data frame above, which has 5 rows and 3 columns, the numerical indexing scheme looks like this:

row	column		
	1	2	3
1	[1,1]	[1,2]	[1,3]
2	[2,1]	[2,2]	[2,3]
3	[3,1]	[3,2]	[3,3]
4	[4,1]	[4,2]	[4,3]
5	[5,1]	[5,2]	[5,3]

If I want the 3rd case of the 2nd variable, what I would type is `garden[3,2]`, and R would print out some output showing that, this element corresponds to the utterance "ee". However, let's hold off from actually doing that for a moment, because there's something slightly counterintuitive about the specifics of what R does under those circumstances (see Section 7.5.4). Instead, let's aim to solve our original problem, which is to pull out two rows (4 and 5) and two columns (1 and 2). This is fairly simple to do, since R allows us to specify multiple rows and multiple columns. So let's try that:

```
> garden[ 4:5, 1:2 ]
   speaker  utterance
case.4 makka-pakka      pip
case.5 makka-pakka      onk
```

Clearly, that's exactly what we asked for: the output here is a data frame containing two variables and two cases. Note that I could have gotten the same answer if I'd used the `c()` function to produce my vectors rather than the `:` operator. That is, the following command is equivalent to the last one:

```
> garden[ c(4,5), c(1,2) ]
```

It's just not as pretty. However, if the columns and rows that you want to keep don't happen to be next to each other in the original data frame, then you might find that you have to resort to using commands like `garden[c(2,4,5), c(1,3)]` to extract them.

A second way to do the same thing is to use the names of the rows and columns. That is, instead of using the row numbers and column numbers, you use the character strings that are used as the labels for the rows and columns. To apply this idea to our `garden` data frame, we would use a command like this:

```
> garden[ c("case.4", "case.5"), c("speaker", "utterance") ]
```

Once again, this produces exactly the same output, so I haven't bothered to show it. Note that, although this version is more annoying to *type* than the previous version, it's a bit easier to *read*, because it's often more meaningful to refer to the elements by their names rather than their numbers. Also note that you don't have to use the same convention for the rows and columns. For instance, I often find that the variable names are meaningful and so I sometimes refer to them by name, whereas the row names are pretty arbitrary so it's easier to refer to them by number. In fact, that's more or less exactly what's happening with the `garden` data frame, so it probably makes more sense to use this as the command:

```
> garden[ 4:5, c("speaker", "utterance") ]
```

Again, the output is identical.

Finally, both the rows and columns can be indexed using logical vectors as well. For example, although I *claimed* earlier that my goal was to extract cases 4 and 5, it's pretty obvious that what I really wanted to do was select the cases where Makka-Pakka is speaking. So what I could have done is create a logical vector that indicates which cases correspond to Makka-Pakka speaking:

```
> is.MP.speaking <- garden$speaker == "makka-pakka"
> is.MP.speaking
[1] FALSE FALSE FALSE  TRUE  TRUE
```

As you can see, the 4th and 5th elements of this vector are `TRUE` while the others are `FALSE`. Now that I've constructed this "indicator" variable, what I can do is use this vector to select the rows that I want to keep:

```
> garden[ is.MP.speaking, c("speaker", "utterance") ]
```

And of course the output is, yet again, the same.

7.5.3 Using square brackets: II. Some elaborations

There are two fairly useful elaborations on this "rows and columns" approach that I should point out. Firstly, what if you want to keep all of the rows, or all of the columns? To do this, all we have to do is leave the corresponding entry blank, but it is crucial to remember to *keep the comma!* For instance, suppose I want to keep all the rows in the `garden` data, but I only want to retain the first two columns. The easiest way to do this is to use a command like this:

```
> garden[ , 1:2 ]
      speaker utterance
case.1  upsy-daisy      pip
case.2  upsy-daisy      pip
case.3  tombliboo       ee
case.4  makka-pakka    pip
case.5  makka-pakka    onk
```

Alternatively, if I want to keep all the columns but only want the last two rows, I use the same trick, but this time I leave the second index blank. So my command becomes:

```
> garden[ 4:5, ]
      speaker utterance line
case.4  makka-pakka    pip    7
case.5  makka-pakka    onk    9
```

The second elaboration I should note is that it's still okay to use negative indexes as a way of telling R to delete certain rows or columns. For instance, if I want to delete the 3rd column, then I use this command:

```
> garden[ , -3 ]
      speaker utterance
case.1  upsy-daisy      pip
case.2  upsy-daisy      pip
case.3  tombliboo       ee
case.4  makka-pakka    pip
case.5  makka-pakka    onk
```

whereas if I want to delete the 3rd row, then I'd use this one:

```
> garden[ -3, ]
      speaker utterance line
case.1  upsy-daisy      pip    1
case.2  upsy-daisy      pip    2
case.4  makka-pakka    pip    7
case.5  makka-pakka    onk    9
```

So that's nice.

7.5.4 Using square brackets: III. Understanding “dropping”

At this point some of you might be wondering why I’ve been so terribly careful to choose my examples in such a way as to ensure that the output always has multiple rows and multiple columns. The reason for this is that I’ve been trying to hide the somewhat curious “dropping” behaviour that R produces when the output only has a single column. I’ll start by showing you what happens, and then I’ll try to explain it. Firstly, let’s have a look at what happens when the output contains only a single *row*:

```
> garden[ 5, ]  
      speaker utterance line  
case.5 makka-pakka      onk      9
```

This is exactly what you’d expect to see: a data frame containing three variables, and only one case per variable. Okay, no problems so far. What happens when you ask for a single *column*? Suppose, for instance, I try this as a command:

```
garden[ , 3 ]
```

Based on everything that I’ve shown you so far, you would be well within your rights to expect to see R produce a data frame containing a single variable (i.e., `line`) and five cases. After all, that *is* what the `subset()` command does in this situation, and it’s pretty consistent with everything else that I’ve shown you so far about how square brackets work. In other words, you should expect to see this:

```
      line  
case.1    1  
case.2    2  
case.3    5  
case.4    7  
case.5    9
```

However, that is emphatically not what happens. What you actually get is this:

```
> garden[ , 3 ]  
[1] 1 2 5 7 9
```

That output is *not a data frame* at all! That’s just an ordinary numeric vector containing 5 elements. What’s going on here is that R has “noticed” that the output that we’ve asked for doesn’t really “need” to be wrapped up in a data frame at all, because it only corresponds to a single variable. So what it does is “drop” the output from a data frame *containing* a single variable, “down” to a simpler output that corresponds to that variable. This behaviour is actually very convenient for day to day usage once you’ve become familiar with it – and I suppose that’s the real reason why R does this – but there’s no escaping the fact that it is *deeply* confusing to novices. It’s especially confusing because the behaviour appears only for a very specific case: (a) it only works for columns and not for rows, because the columns correspond to variables and the rows do not, and (b) it only applies to the “rows and columns” version of the square brackets, and not to the `subset()` function,¹¹ or to the “just columns” use of the square brackets (next section). As I say, it’s very confusing when you’re just starting out. For what it’s worth, you can suppress this behaviour if you want, by setting `drop = FALSE` when you construct your bracketed expression. That is, you could do something like this:

```
> garden[ , 3, drop = FALSE ]  
      line
```

¹¹Actually, you can make the `subset()` function behave this way by using the optional `drop` argument, but by default `subset()` does not drop, which is probably more sensible and more intuitive to novice users.

```

case.1    1
case.2    2
case.3    5
case.4    7
case.5    9

```

I suppose that helps a little bit, in that it gives you some control over the dropping behaviour, but I'm not sure it helps to make things any easier to understand. Anyway, that's the "dropping" special case. Fun, isn't it?

7.5.5 Using square brackets: IV. Columns only

As if the weird "dropping" behaviour wasn't annoying enough, R actually provides a completely different way of using square brackets to index a data frame. Specifically, if you *only* give a single index, R will assume you want the corresponding columns, not the rows. Do not be fooled by the fact that this second method also uses square brackets: it behaves differently to the "rows and columns" method that I've discussed in the last few sections. Again, what I'll do is show you *what* happens first, and then I'll try to explain *why* it happens afterwards. To that end, let's start with the following command:

```

> garden[ 1:2 ]
      speaker utterance
case.1  upsy-daisy      pip
case.2  upsy-daisy      pip
case.3  tombliboo       ee
case.4  makka-pakka     pip
case.5  makka-pakka     onk

```

As you can see, the output gives me the first two columns, much as if I'd typed `garden[,1:2]`. It doesn't give me the first two rows, which is what I'd have gotten if I'd used a command like `garden[1:2,]`. Not only that, if I ask for a *single* column, R does not drop the output:

```

> garden[3]
      line
case.1    1
case.2    2
case.3    5
case.4    7
case.5    9

```

As I said earlier, the *only* case where dropping occurs by default is when you use the "row and columns" version of the square brackets, and the output happens to correspond to a single column. However, if you really want to force R to drop the output, you can do so using the "double brackets" notation:

```

> garden[[3]]
[1] 1 2 5 7 9

```

Note that R will only allow you to ask for one column at a time using the double brackets. If you try to ask for multiple columns in this way, you get completely different behaviour,¹² which may or may not produce an error, but definitely won't give you the output you're expecting. The only reason I'm mentioning it at all is that you might run into double brackets when doing further reading, and a lot

¹²Specifically, recursive indexing, a handy tool in some contexts but not something that I want to discuss here.

of books don't explicitly point out the difference between `[` and `[[`. However, I promise that I won't be using `[[` anywhere else in this book.

Okay, for those few readers that have persevered with this section long enough to get here without having set fire to the book, I should explain *why* R has these two different systems for subsetting a data frame (i.e., "row and column" versus "just columns"), and why they behave so differently to each other. I'm not 100% sure about this since I'm still reading through some of the old references that describe the early development of R, but I think the answer relates to the fact that data frames are actually a very strange hybrid of two different kinds of thing. At a low level, a data frame is a list (Section 4.9). I can demonstrate this to you by overriding the normal `print()` function¹³ and forcing R to print out the `garden` data frame using the default print method rather than the special one that is defined only for data frames. Here's what we get:

```
> print.default( garden )
$speaker
[1] upsy-daisy  upsy-daisy  tombliboo  makka-pakka makka-pakka
Levels: makka-pakka tombliboo upsy-daisy

$utterance
[1] pip pip ee  pip onk
Levels: ee onk oo pip

$line
[1] 1 2 5 7 9

attr("class")
[1] "data.frame"
```

Apart from the weird part of the output right at the bottom, this is *identical* to the print out that you get when you print out a list (see Section 4.9). In other words, a data frame is a list. View from this "list based" perspective, it's clear what `garden[1]` is: it's the first variable stored in the list, namely `speaker`. In other words, when you use the "just columns" way of indexing a data frame, using only a single index, R assumes that you're thinking about the data frame as if it were a *list of variables*. In fact, when you use the `$` operator you're taking advantage of the fact that the data frame is secretly a list.

However, a data frame is more than just a list. It's a very special kind of list where all the variables are of the same length, and the first element in each variable happens to correspond to the first "case" in the data set. That's why no-one ever wants to see a data frame printed out in the default "list-like" way that I've shown in the extract above. In terms of the deeper *meaning* behind what a data frame is used for, a data frame really does have this rectangular shape to it:

```
> print( garden )
      speaker utterance line
case.1  upsy-daisy      pip     1
case.2  upsy-daisy      pip     2
case.3   tombliboo      ee      5
case.4  makka-pakka     pip     7
case.5  makka-pakka    onk     9
```

Because of the fact that a data frame is basically a table of data, R provides a second "row and column" method for interacting with the data frame (see Section 7.11.1 for a related example). This method makes much more sense in terms of the high-level *table of data* interpretation of what a data frame is, and so for the most part it's this method that people tend to prefer. In fact, throughout the rest of the book I will

¹³Remember, `print()` is generic: see Section 4.11.

be sticking to the “row and column” approach (though I will use `$` a lot), and never again referring to the “just columns” approach. However, it does get used a lot in practice, so I think it’s important that this book explain what’s going on.

And now let us never speak of this again.

7.6

Sorting, flipping and merging data

In this section I discuss a few useful operations that I feel are loosely related to one another: sorting a vector, sorting a data frame, binding two or more vectors together into a data frame (or matrix), and flipping a data frame (or matrix) on its side. They’re all fairly straightforward tasks, at least in comparison to some of the more obnoxious data handling problems that turn up in real life.

7.6.1 Sorting a numeric or character vector

One thing that you often want to do is sort a variable. If it’s a numeric variable you might want to sort in increasing or decreasing order. If it’s a character vector you might want to sort alphabetically, etc. The `sort()` function provides this capability.

```
> numbers <- c(2,4,3)
> sort( x = numbers )
[1] 2 3 4
```

You can ask for R to sort in decreasing order rather than increasing:

```
> sort( x = numbers, decreasing = TRUE )
[1] 4 3 2
```

And you can ask it to sort text data in alphabetical order:

```
> text <- c("aardvark", "zebra", "swing")
> sort( text )
[1] "aardvark" "swing"     "zebra"
```

That’s pretty straightforward. That being said, it’s important to note that I’m glossing over something here. When you apply `sort()` to a character vector it doesn’t strictly sort into alphabetical order. R actually has a slightly different notion of how characters are ordered (see Section 7.8.5 and Table 7.3), which is more closely related to how computers store text data than to how letters are ordered in the alphabet. However, that’s a topic we’ll discuss later. For now, the only thing I should note is that the `sort()` function doesn’t alter the original variable. Rather, it creates a new, sorted variable as the output. So if I inspect my original `text` variable:

```
> text
[1] "aardvark" "zebra"     "swing"
```

I can see that it has remained unchanged.

7.6.2 Sorting a factor

You can also sort factors, but the story here is slightly more subtle because there's two different ways you can sort a factor: alphabetically (by label) or by factor level. The `sort()` function uses the latter. To illustrate, let's look at the two different examples. First, let's create a factor in the usual way:

```
> fac <- factor( text )
> fac
[1] aardvark zebra      swing
Levels: aardvark swing zebra
```

Now let's sort it:

```
> sort(fac)
[1] aardvark swing      zebra
Levels: aardvark swing zebra
```

This *looks* like it's sorted things into alphabetical order, but that's only because the factor levels themselves happen to be alphabetically ordered. Suppose I deliberately define the factor levels in a non-alphabetical order:

```
> fac <- factor( text, levels = c("zebra", "swing", "aardvark") )
> fac
[1] aardvark zebra      swing
Levels: zebra swing aardvark
```

Now what happens when we try to sort `fac` this time? The answer:

```
> sort(fac)
[1] zebra      swing      aardvark
Levels: zebra swing aardvark
```

It sorts the data into the numerical order implied by the factor levels, not the alphabetical order implied by the labels attached to those levels. Normally you never notice the distinction, because by default the factor levels are assigned in alphabetical order, but it's important to know the difference:

7.6.3 Sorting a data frame

The `sort()` function doesn't work properly with data frames. If you want to sort a data frame the standard advice that you'll find online is to use the `order()` function (not described in this book) to determine what order the rows should be sorted, and then use square brackets to do the shuffling. There's nothing inherently wrong with this advice, I just find it tedious. To that end, the `lsr` package includes a function called `sortFrame()` that you can use to do the sorting. The first argument to the function is named `(x)`, and should correspond to the data frame that you want sorted. After that, all you do is type a list of the names of the variables that you want to use to do the sorting. For instance, if I type this:

```
> sortFrame( garden, speaker, line )
   speaker utterance line
case.4 makka-pakka      pip    7
case.5 makka-pakka      onk    9
case.3 tombliboo        ee     5
case.1 upsy-daisy       pip    1
case.2 upsy-daisy       pip    2
```

what R does is first sort by `speaker` (factor level order). Any ties (i.e., data from the same speaker) are then sorted in order of `line` (increasing numerical order). You can use the minus sign to indicate that numerical variables should be sorted in reverse order:

```
> sortFrame( garden, speaker, -line)
      speaker utterance line
case.5 makka-pakka    onk     9
case.4 makka-pakka    pip     7
case.3 tombliboo      ee      5
case.2 upsy-daisy     pip     2
case.1 upsy-daisy     pip     1
```

As of the current writing, the `sortFrame()` function is under development. I've started introducing functionality to allow you to use the `-` sign to non-numeric variables or to make a distinction between sorting factors alphabetically or by factor level. The idea is that you should be able to type in something like this:

```
> sortFrame( garden, -speaker)
```

and have the output correspond to a sort of the `garden` data frame in *reverse* alphabetical order (or reverse factor level order) of `speaker`. As things stand right now, this will actually work, and it will produce sensible output:

```
> sortFrame( garden, -speaker)
      speaker utterance line
case.1 upsy-daisy     pip     1
case.2 upsy-daisy     pip     2
case.3 tombliboo      ee      5
case.4 makka-pakka    pip     7
case.5 makka-pakka    onk     9
```

However, I'm not completely convinced that I've set this up in the ideal fashion, so this may change a little bit in the future.

7.6.4 Binding vectors together

A not-uncommon task that you might find yourself needing to undertake is to combine several vectors. For instance, let's suppose we have the following two numeric vectors:

```
> cake.1 <- c(100, 80, 0, 0, 0)
> cake.2 <- c(100, 100, 90, 30, 10)
```

The numbers here might represent the amount of each of the two cakes that are left at five different time points. Apparently the first cake is tastier, since that one gets devoured faster. We've already seen one method for combining these vectors: we could use the `data.frame()` function to convert them into a data frame with two variables, like so:

```
> cake.df <- data.frame( cake.1, cake.2 )
> cake.df
  cake.1 cake.2
1     100     100
2      80     100
3       0      90
4       0      30
5       0      10
```

Two other methods that I want to briefly refer to are the `rbind()` and `cbind()` functions, which will convert the vectors into a matrix. I'll discuss matrices properly in Section 7.11.1 but the details don't matter too much for our current purposes. The `cbind()` function ("column bind") produces a very similar looking output to the data frame example:

```
> cake.mat1 <- cbind( cake.1, cake.2 )
> cake.mat1
      cake.1 cake.2
[1,]    100    100
[2,]     80    100
[3,]      0     90
[4,]      0     30
[5,]      0     10
```

but nevertheless it's important to keep in mind that `cake.mat1` is a matrix rather than a data frame, and so has a few differences from the `cake.df` variable. The `rbind()` function ("row bind") produces a somewhat different output: it binds the vectors together row-wise rather than column-wise, so the output now looks like this:

```
> cake.mat2 <- rbind( cake.1, cake.2 )
> cake.mat2
      [,1] [,2] [,3] [,4] [,5]
cake.1  100   80    0    0    0
cake.2   100   100   90   30   10
```

You can add names to a matrix by using the `rownames()` and `colnames()` functions, and I should also point out that there's a fancier function in R called `merge()` that supports more complicated "database like" merging of vectors and data frames, but I won't go into details here.

7.6.5 Binding multiple copies of the same vector together

It is sometimes very useful to bind together multiple copies of the same vector. You could do this using the `rbind` and `cbind` functions, using commands like this one

```
> fibonacci <- c( 1,1,2,3,5,8 )
> rbind( fibonacci, fibonacci, fibonacci )
      [,1] [,2] [,3] [,4] [,5] [,6]
fibonacci    1    1    2    3    5    8
fibonacci    1    1    2    3    5    8
fibonacci    1    1    2    3    5    8
```

but that can be pretty annoying, especially if you need lots of copies. To make this a little easier, the `lqr` package has two additional functions `rowCopy` and `colCopy` that do the same job, but all you have to do is specify the number of copies that you want, instead of typing the name in over and over again. The two arguments you need to specify are `x`, the vector to be copied, and `times`, indicating how many copies should be created:¹⁴

```
> rowCopy( x = fibonacci, times = 3 )
      [,1] [,2] [,3] [,4] [,5] [,6]
```

¹⁴Note for advanced users: both of these functions are just wrappers to the `matrix()` function, which is pretty flexible in terms of the ability to convert vectors into matrices. Also, while I'm on this topic, I'll briefly mention the fact that if you're a Matlab user and looking for an equivalent of Matlab's `repmat()` function, I'd suggest checking out the `matlab` package which contains R versions of a lot of handy Matlab functions.

```
[1,] 1 1 2 3 5 8
[2,] 1 1 2 3 5 8
[3,] 1 1 2 3 5 8
```

Of course, in practice you don't need to name the arguments all the time. For instance, here's an example using the `colCopy()` function with the argument names omitted:

```
> colCopy( fibonacci, 3 )
[,1] [,2] [,3]
[1,] 1 1 1
[2,] 1 1 1
[3,] 2 2 2
[4,] 3 3 3
[5,] 5 5 5
[6,] 8 8 8
```

7.6.6 Transposing a matrix or data frame

One of the main reasons that I wanted to discuss the `rbind()` and `cbind()` functions in the same section as the `data.frame()` function is that it immediately raises the question of how to "flip" or `transpose` a matrix or data frame. Notice that in the last section I was able to produce two different matrices, `cake.mat1` and `cake.mat2` that were basically mirror images of one another. A natural question to ask is whether you can directly transform one into another. The transpose function `t()` allows us to do this in a straightforward fashion. To start with, I'll show you how to transpose a matrix, and then I'll move onto talk about data frames. Firstly, let's load a matrix I prepared earlier, from the `cakes.Rdata` file:

```
> load( "cakes.Rdata" )
> cakes
   cake.1 cake.2 cake.3 cake.4
time.1    100    100    100    100
time.2     80    100     20    100
time.3      0     90     20    100
time.4      0     30     20    100
time.5      0     10     20    100
```

And just to make sure you believe me that this is actually a matrix:

```
> class( cakes )
[1] "matrix"
```

Okay, now let's transpose the matrix:

```
> cakes.flipped <- t( cakes )
> cakes.flipped
   time.1 time.2 time.3 time.4 time.5
cake.1    100     80      0      0      0
cake.2    100    100     90     30     10
cake.3    100     20     20     20     20
cake.4    100    100    100    100    100
```

The output here is still a matrix:

```
> class( cakes.flipped )
[1] "matrix"
```

At this point you should have two questions: (1) how do we do the same thing for data frames? and (2) why should we care about this? Let's start with the how question. First, I should note that you can transpose a data frame just fine using the `t()` function, but that has the slightly awkward consequence of converting the output from a data frame to a matrix, which isn't usually what you want. It's quite easy to convert the output back again, of course,¹⁵ but I hate typing two commands when I can do it with one. To that end, the `lsr` package has a simple "convenience" function called `tFrame()` which does exactly the same thing as `t()` but converts the output to a data frame for you. To illustrate this, let's transpose the `itng` data frame that we used earlier. Here's the original data frame:

```
> itng
      speaker utterance
 1  upsy-daisy      pip
 2  upsy-daisy      pip
 3  upsy-daisy     onk
 4  upsy-daisy     onk
 5  tombliboo      ee
 6  tombliboo      oo
 7 makka-pakka      pip
 8 makka-pakka      pip
 9 makka-pakka     onk
10 makka-pakka     onk
```

and here's what happens when you transpose it using `tFrame()`:

```
> tFrame( itng )
      V1      V2      V3      V4      V5
speaker  upsy-daisy upsy-daisy upsy-daisy upsy-daisy tombliboo
utterance    pip        pip      onk      onk      ee
      V6      V7      V8      V9      V10
speaker   tombliboo makka-pakka makka-pakka makka-pakka makka-pakka
utterance     oo        pip      pip      onk      onk
```

An important point to recognise is that transposing a data frame is not always a sensible thing to do: in fact, I'd go so far as to argue that it's usually *not* sensible. It depends a lot on whether the "cases" from your original data frame would make sense as variables, and to think of each of your original "variables" as cases. I think that's emphatically *not* true for our `itng` data frame, so I wouldn't advise doing it in this situation.

That being said, sometimes it really is true. For instance, had we originally stored our `cakes` variable as a data frame instead of a matrix, then it would absolutely be sensible to flip the data frame!¹⁶ There are some situations where it is useful to flip your data frame, so it's nice to know that you can do it. Indeed, that's the main reason why I have spent so much time talking about this topic. A lot of statistical tools make the assumption that the rows of your data frame (or matrix) correspond to observations, and the columns correspond to the variables. That's not unreasonable, of course, since that is a pretty standard convention. However, think about our `cakes` example here. This is a situation where you might want do an analysis of the different cakes (i.e. cakes as variables, time points as cases), but equally you might want to do an analysis where you think of the times as being the things of interest (i.e., times as variables, cakes as cases). If so, then it's useful to know how to flip a matrix or data frame around.

¹⁵The function you need for that is called `as.data.frame()`.

¹⁶In truth, I suspect that most of the cases when you can sensibly flip a data frame occur when all of the original variables are measurements of the same type (e.g., all variables are response times), and if so you could easily have chosen to encode your data as a matrix instead of as a data frame. But since people do sometimes prefer to work with data frames, I've written the `tFrame()` function for the sake of convenience. I don't really think it's something that is needed very often.

Reshaping a data frame

One of the most annoying tasks that you need to undertake on a regular basis is that of reshaping a data frame. Framed in the most general way, reshaping the data means taking the data in whatever format it's given to you, and converting it to the format you need it. Of course, if we're going to characterise the problem that broadly, then about half of this chapter can probably be thought of as a kind of reshaping. So we're going to have to narrow things down a little bit. To that end, I'll talk about a few different tools that you can use for a few different tasks. In particular, I'll discuss a couple of easy to use (but limited) functions that I've included in the `lsr` package. In future versions of the book I plan to expand this discussion to include some of the more powerful tools that are available in R, but I haven't had the time to do so yet.

7.7.1 Long form and wide form data

The most common format in which you might obtain data is as a “case by variable” layout, commonly known as the **wide form** of the data.

```
> load("repeated.Rdata")
> who()
-- Name --  -- Class --  -- Size --
choice      data.frame    4 x 10
drugs       data.frame   10 x 8
```

To get a sense of what I'm talking about, consider an experiment in which we are interested in the different effects that alcohol and caffeine have on people's working memory capacity (WMC) and reaction times (RT). We recruit 10 participants, and measure their WMC and RT under three different conditions: a “no drug” condition, in which they are not under the influence of either caffeine or alcohol, a “caffeine” condition, in which they are under the influence of caffeine, and an “alcohol” condition, in which... well, you can probably guess. Ideally, I suppose, there would be a fourth condition in which both drugs are administered, but for the sake of simplicity let's ignore that. The `drugs` data frame gives you a sense of what kind of data you might observe in an experiment like this:

```
> drugs
  id gender WMC_alcohol WMC_caffeine WMC_no.drug RT_alcohol RT_caffeine RT_no.drug
  1  1 female     3.7        3.7      3.9       488      236      371
  2  2 female     6.4        7.3      7.9       607      376      349
  3  3 female     4.6        7.4      7.3       643      226      412
  4  4 male       6.4        7.8      8.2       684      206      252
  5  5 female     4.9        5.2      7.0       593      262      439
  6  6 male       5.4        6.6      7.2       492      230      464
  7  7 male       7.9        7.9      8.9       690      259      327
  8  8 male       4.1        5.9      4.5       486      230      305
  9  9 female     5.2        6.2      7.2       686      273      327
 10 10 female     6.2        7.4      7.8       645      240      498
```

This is a data set in “wide form”, in which each participant corresponds to a single row. We have two variables that are characteristics of the subject (i.e., their `id` number and their `gender`) and six variables that refer to one of the two measured variables (WMC or RT) in one of the three testing conditions

(alcohol, caffeine or no drug). Because all of the testing conditions (i.e., the three drug types) are applied to all participants, drug type is an example of a **within-subject factor**.

7.7.2 Reshaping data using `wideToLong()`

The “wide form” of this data set is useful for some situations: it is often very useful to have each row correspond to a single subject. However, it is not the only way in which you might want to organise this data. For instance, you might want to have a separate row for each “testing occasion”. That is, “participant 1 under the influence of alcohol” would be one row, and “participant 1 under the influence of caffeine” would be another row. This way of organising the data is generally referred to as the **long form** of the data. It’s not too difficult to switch between wide and long form, and I’ll explain how it works in a moment; for now, let’s just have a look at what the long form of this data set looks like:

```
> drugs.2 <- wideToLong( data = drugs, within = "drug" )
> drugs.2
   id gender     drug WMC  RT
 1  1 female alcohol 3.7 488
 2  2 female alcohol 6.4 607
 3  3 female alcohol 4.6 643
 4  4 male   alcohol 6.4 684
 5  5 female alcohol 4.9 593
 6  6 male   alcohol 5.4 492
 7  7 male   alcohol 7.9 690
 8  8 male   alcohol 4.1 486
 9  9 female alcohol 5.2 686
10 10 female alcohol 6.2 645
11  1 female caffeine 3.7 236
12  2 female caffeine 7.3 376
```

BLAH BLAH BLAH

The `drugs.2` data frame that we just created has 30 rows: each of the 10 participants appears in three separate rows, one corresponding to each of the three testing conditions. And instead of having a variable like `WMC_caffeine` that indicates that we were measuring “WMC” in the “caffeine” condition, this information is now recorded in two separate variables, one called `drug` and another called `WMC`. Obviously, the long and wide forms of the data contain the same information, but they represent quite different ways of organising that information. Sometimes you find yourself needing to analyse data in wide form, and sometimes you find that you need long form. So it’s really useful to know how to switch between the two.

In the example I gave above, I used a function called `wideToLong()` to do the transformation. The `wideToLong()` function is part of the `lsr` package. The key to understanding this function is that it relies on the *variable names* to do all the work. Notice that the variable names in the `drugs` data frame follow a very clear scheme. Whenever you have a variable with a name like `WMC_caffeine` you know that the variable being measured is “WMC”, and that the specific condition in which it is being measured is the “caffeine” condition. Similarly, you know that `RT_no.drug` refers to the “RT” variable measured in the “no drug” condition. The measured variable comes first (e.g., `WMC`), followed by a separator character (in this case the separator is an underscore, `_`), and then the name of the condition in which it is being measured (e.g., `caffeine`). There are two different prefixes (i.e, the strings before the separator, `WMC`, `RT`) which means that there are two separate variables being measured. There are three different suffixes (i.e., the strings after the separator, `caffeine`, `alcohol`, `no.drug`) meaning that there are three different levels of the within-subject factor. Finally, notice that the separator string (i.e., `_`) does not appear anywhere in two of the variables (`id`, `gender`), indicating that these are **between-subject** variables, namely variables that do not vary within participant (e.g., a person’s `gender` is the same regardless of whether they’re

under the influence of alcohol, caffeine etc).

Because of the fact that the variable naming scheme here is so informative, it's quite possible to reshape the data frame without any additional input from the user. For example, in this particular case, you could just type the following:

```
> wideToLong( drugs )
   id gender  within WMC  RT
1  1 female  alcohol 3.7 488
2  2 female  alcohol 6.4 607
3  3 female  alcohol 4.6 643
4  4 male   alcohol 6.4 684

BLAH BLAH BLAH
```

This is pretty good, actually. The only think it has gotten wrong here is that it doesn't know what name to assign to the within-subject factor, so instead of calling it something sensible like `drug`, it has use the unimaginative name `within`. If you want to ensure that the `wideToLong()` function applies a sensible name, you have to specify the `within` argument, which is just a character string that specifies the name of the within-subject factor. So when I used this command earlier,

```
> drugs.2 <- wideToLong( data = drugs, within = "drug" )
```

all I was doing was telling R to use `drug` as the name of the within subject factor.

Now, as I was hinting earlier, the `wideToLong()` function is very inflexible. It *requires* that the variable names all follow this naming scheme that I outlined earlier. If you don't follow this naming scheme it won't work.¹⁷ The only flexibility that I've included here is that you can change the separator character by specifying the `sep` argument. For instance, if you were using variable names of the form `WMC/caffeine`, for instance, you could specify that `sep="/"`, using a command like this

```
> drugs.2 <- wideToLong( data = drugs, within = "drug", sep = "/" )
```

and it would still work.

7.7.3 Reshaping data using `longToWide()`

To convert data from long form to wide form, the `lsr` package also includes a function called `longToWide()`. Recall from earlier that the long form of the data (i.e., the `drugs.2` data frame) contains variables named `id`, `gender`, `drug`, `WMC` and `RT`. In order to convert from long form to wide form, all you need to do is indicate which of these variables are measured separately for each condition (i.e., `WMC` and `RT`), and which variable is the within-subject factor that specifies the condition (i.e., `drug`). You do this via a two-sided formula, in which the measured variables are on the left hand side, and the within-subject factor is on the right hand side. In this case, the formula would be `WMC + RT ~ drug`. So the command that we would use might look like this:

```
> longToWide( data=drugs.2, formula= WMC+RT ~ drug )
   id gender WMC_alcohol RT_alcohol WMC_caffeine RT_caffeine WMC_no.drug RT_no.drug
1  1 female      3.7     488      3.7     236      3.9     371
2  2 female      6.4     607      7.3     376      7.9     349
```

¹⁷This limitation is deliberate, by the way: if you're getting to the point where you want to do something more complicated, you should probably start learning how to use `reshape()`, `cast()` and `melt()` or some of other the more advanced tools. The `wideToLong()` and `longToWide()` functions are included only to help you out when you're first starting to use R.

3	3	female	4.6	643	7.4	226	7.3	412
4	4	male	6.4	684	7.8	206	8.2	252
5	5	female	4.9	593	5.2	262	7.0	439
6	6	male	5.4	492	6.6	230	7.2	464
7	7	male	7.9	690	7.9	259	8.9	327
8	8	male	4.1	486	5.9	230	4.5	305
9	9	female	5.2	686	6.2	273	7.2	327
10	10	female	6.2	645	7.4	240	7.8	498

or, if we chose to omit argument names, we could simplify it to this:

```
> longToWide( drugs.2, WMC+RT ~ drug )
```

Note that, just like the `wideToLong()` function, the `longToWide()` function allows you to override the default separator character. For instance, if the command I used had been

```
> longToWide( drugs.2, WMC+RT ~ drug, sep="/" )
```

the output would contain variables with names like `RT/alcohol` instead of `RT_alcohol`.

7.7.4 Reshaping with multiple within-subject factors

As I mentioned above, the `wideToLong()` and `longToWide()` functions are quite limited in terms of what they can do. However, they do handle a broader range of situations than the one outlined above. Consider the following, fairly simple psychological experiment. I'm interested in the effects of practice on some simple decision making problem. It doesn't really matter what the problem is, other than to note that I'm interested in two distinct outcome variables. Firstly, I care about people's accuracy, measured by the proportion of decisions that people make correctly, denoted PC. Secondly, I care about people's speed, measured by the mean response time taken to make those decisions, denoted MRT. That's standard in psychological experiments: the speed-accuracy trade-off is pretty ubiquitous, so we generally need to care about both variables.

To look at the effects of practice over the long term, I test each participant on two days, `day1` and `day2`, where for the sake of argument I'll assume that `day1` and `day2` are about a week apart. To look at the effects of practice over the short term, the testing during each day is broken into two "blocks", `block1` and `block2`, which are about 20 minutes apart. This isn't the world's most complicated experiment, but it's still a fair bit more complicated than the last one. This time around we have two within-subject factors (i.e., `day` and `block`) and we have two measured variables for each condition (i.e., `PC` and `MRT`). The `choice` data frame shows what the wide form of this kind of data might look like:

```
> choice
```

	id	gender	MRT/block1/day1	MRT/block1/day2	MRT/block2/day1	MRT/block2/day2
1	1	male	415	400	455	450
2	2	male	500	490	532	518
3	3	female	478	468	499	474
4	4	female	550	502	602	588

	PC/block1/day1	PC/block1/day2	PC/block2/day1	PC/block2/day2
1	79	88	82	93
2	83	92	86	97
3	91	98	90	100
4	75	89	78	95

Notice that this time around we have variable names of the form `MRT/block1/day2`. As before, the first part of the name refers to the measured variable (response time), but there are now two suffixes, one indicating that the testing took place in block 1, and the other indicating that it took place on day 2. And just to complicate matters, it uses `/` as the separator character rather than `_`. Even so, reshaping this data set is pretty easy. The command to do it is,

```
> choice.2 <- wideToLong( choice, within=c("block","day"), sep="/" )
```

which is pretty much the exact same command we used last time. The only difference here is that, because there are two within-subject factors, the `within` argument is a vector that contains two names. When we look at the long form data frame that this creates, we get this:

```
> choice.2
   id gender MRT  PC  block  day
1   1   male  415  79 block1 day1
2   2   male  500  83 block1 day1
3   3 female  478  91 block1 day1
4   4 female  550  75 block1 day1
5   1   male  400  88 block1 day2
6   2   male  490  92 block1 day2

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15  3 female  474 100 block2 day2
16  4 female  588  95 block2 day2
```

In this long form data frame we have two between-subject variables (`id` and `gender`), two variables that define our within-subject manipulations (`block` and `day`), and two more contain the measurements we took (`MRT` and `PC`).

To convert this back to wide form is equally straightforward. We use the `longToWide()` function, but this time around we need to alter the formula in order to tell it that we have two within-subject factors. The command is now

```
> longToWide( choice.2, MRT+PC ~ block+day, sep="/" )
```

and this produces a wide form data set containing the same variables as the original `choice` data frame.

7.7.5 What other options are there?

The advantage to the approach described in the previous section is that it solves a quite specific problem (but a commonly encountered one) with a minimum of fuss. The disadvantage is that the tools are quite limited in scope. They allow you to switch your data back and forth between two different formats that are very common in everyday data analysis. However, there are a number of other tools that you can use if need be. Just within the core packages distributed with R there is the `reshape()` function, as well as the `stack()` and `unstack()` functions, all of which can be useful under certain circumstances. And there are of course thousands of packages on CRAN that you can use to help you with different tasks. One popular package for this purpose is the `reshape` package, written by Hadley Wickham (see [Wickham, 2007](#), for details). There are two key functions in this package, called `melt()` and `cast()` that are pretty useful for solving a lot of reshaping problems. In a future version of this book I intend to

discuss `melt()` and `cast()` in a fair amount of detail.

7.8

Working with text

Sometimes your data set is quite text heavy. This can be for a lot of different reasons. Maybe the raw data are actually taken from text sources (e.g., newspaper articles), or maybe your data set contains a lot of free responses to survey questions, in which people can write whatever text they like in response to some query. Or maybe you just need to rejig some of the text used to describe nominal scale variables. Regardless of what the reason is, you'll probably want to know a little bit about how to handle text in R. Some things you already know how to do: I've discussed the use of `nchar()` to calculate the number of characters in a string (Section 3.7.1), and a lot of the general purpose tools that I've discussed elsewhere (e.g., the `==` operator) have been applied to text data as well as to numeric data. However, because text data is quite rich, and generally not as well structured as numeric data, R provides a lot of additional tools that are quite specific to text. In this section I discuss only those tools that come as part of the base packages, but there are other possibilities out there: the `stringr` package provides a powerful alternative that is a lot more coherent than the basic tools, and is well worth looking into.

7.8.1 Shortening a string

The first task I want to talk about is how to shorten a character string. For example, suppose that I have a vector that contains the names of several different animals:

```
> animals <- c( "cat", "dog", "kangaroo", "whale" )
```

It might be useful in some contexts to extract the first three letters of each word. This is often useful when annotating figures, or when creating variable labels: it's often very inconvenient to use the full name, so you want to shorten it to a short code for space reasons. The `strtrim()` function can be used for this purpose. It has two arguments: `x` is a vector containing the text to be shortened and `width` specifies the number of characters to keep. When applied to the `animals` data, here's what we get:

```
> strtrim( x = animals, width = 3 )
[1] "cat"  "dog"  "kan"  "wha"
```

Note that the only thing that `strtrim()` does is chop off excess characters at the end of a string. It doesn't insert any whitespace characters to fill them out if the original string is shorter than the `width` argument. For example, if I trim the `animals` data to 4 characters, here's what I get:

```
> strtrim( x = animals, width = 4 )
[1] "cat"  "dog"  "kang" "whal"
```

The `"cat"` and `"dog"` strings still only use 3 characters. Okay, but what if you don't want to start from the first letter? Suppose, for instance, I only wanted to keep the second and third letter of each word. That doesn't happen quite as often, but there are some situations where you need to do something like that. If that does happen, then the function you need is `substr()`, in which you specify a `start` point and a `stop` point instead of specifying the width. For instance, to keep only the 2nd and 3rd letters of the various `animals`, I can do the following:

```
> substr( x = animals, start = 2, stop = 3 )
[1] "at"   "og"   "an"   "ha"
```

7.8.2 Pasting strings together

Much more commonly, you will need either to glue several character strings together or to pull them apart. To glue several strings together, the `paste()` function is very useful. There are three arguments to the `paste()` function:

- ... As usual, the dots “match” up against any number of inputs. In this case, the inputs should be the various different strings you want to paste together.
- `sep`. This argument should be a string, indicating what characters R should use as separators, in order to keep each of the original strings separate from each other in the pasted output. By default the value is a single space, `sep = " "`. This is made a little clearer when we look at the examples.
- `collapse`. This is an argument indicating whether the `paste()` function should interpret vector inputs as things to be collapsed, or whether a vector of inputs should be converted into a vector of outputs. The default value is `collapse = NULL` which is interpreted as meaning that vectors should not be collapsed. If you want to collapse vectors into a single string, then you should specify a value for `collapse`. Specifically, the value of `collapse` should correspond to the separator character that you want to use for the collapsed inputs. Again, see the examples below for more details.

That probably doesn’t make much sense yet, so let’s start with a simple example. First, let’s try to paste two words together, like this:

```
> paste( "hello", "world" )
[1] "hello world"
```

Notice that R has inserted a space between the `"hello"` and `"world"`. Suppose that’s not what I wanted. Instead, I might want to use `.` as the separator character, or to use no separator at all. To do either of those, I would need to specify `sep = ".."` or `sep = ""`.¹⁸ For instance:

```
> paste( "hello", "world", sep = ".")
[1] "hello.world"
```

Now let’s consider a slightly more complicated example. Suppose I have two vectors that I want to `paste()` together. Let’s say something like this:

```
> hw <- c( "hello", "world" )
> ng <- c( "nasty", "government" )
```

And suppose I want to paste these together. However, if you think about it, this statement is kind of ambiguous. It could mean that I want to do an “element wise” paste, in which all of the first elements get pasted together (`"hello nasty"`) and all the second elements get pasted together (`"world government"`). Or, alternatively, I might intend to collapse everything into one big string (`"hello nasty world government"`). By default, the `paste()` function assumes that you want to do an element-wise paste:

```
> paste( hw, ng )
[1] "hello nasty"      "world government"
```

However, there’s nothing stopping you from overriding this default. All you have to do is specify a value for the `collapse` argument, and R will chuck everything into one dirty big string. To give you a sense of exactly how this works, what I’ll do in this next example is specify *different* values for `sep` and `collapse`:

¹⁸To be honest, it does bother me a little that the default value of `sep` is a space. Normally when I want to paste strings together I don’t want any separator character, so I’d prefer it if the default were `sep=""`. To that end, it’s worth noting that there’s also a `paste0()` function, which is identical to `paste()` except that it always assumes that `sep=""`. Type `?paste` for more information about this.

```
> paste( hw, ng, sep = ".", collapse = "::::")
[1] "hello.nasty:::world.government"
```

7.8.3 Splitting strings

At other times you have the opposite problem to the one in the last section: you have a whole lot of text bundled together into a single string that needs to be pulled apart and stored as several different variables. For instance, the data set that you get sent might include a single variable containing someone's full name, and you need to separate it into first names and last names. To do this in R you can use the `strsplit()` function, and for the sake of argument, let's assume that the string you want to split up is the following string:

```
> monkey <- "It was the best of times. It was the blurst of times."
```

To use the `strsplit()` function to break this apart, there are three arguments that you need to pay particular attention to:

- `x`. A vector of character strings containing the data that you want to split.
- `split`. Depending on the value of the `fixed` argument, this is either a fixed string that specifies a delimiter, or a regular expression that matches against one or more possible delimiters. If you don't know what regular expressions are (probably most readers of this book), don't use this option. Just specify a separator string, just like you would for the `paste()` function.
- `fixed`. Set `fixed = TRUE` if you want to use a fixed delimiter. As noted above, unless you understand regular expressions this is definitely what you want. However, the default value is `fixed = FALSE`, so you have to set it explicitly.

Let's look at a simple example:

```
> monkey.1 <- strsplit( x = monkey, split = " ", fixed = TRUE )
> monkey.1
[[1]]
[1] "It"      "was"     "the"     "best"    "of"      "times." "It"      "was"
[9] "the"     "blurst"  "of"      "times."
```

One thing to note in passing is that the output here is a list (you can tell from the `[[1]]` part of the output), whose first and only element is a character vector. This is useful in a lot of ways, since it means that you can input a character vector for `x` and then have the `strsplit()` function split all of them, but it's kind of annoying when you only have a single input. To that end, it's useful to know that you can `unlist()` the output:

```
> unlist( monkey.1 )
[1] "It"      "was"     "the"     "best"    "of"      "times." "It"      "was"
[9] "the"     "blurst"  "of"      "times."
```

To understand why it's important to remember to use the `fixed = TRUE` argument, suppose we wanted to split this into two separate sentences. That is, we want to use `split = "."` as our delimiter string. As long as we tell R to remember to treat this as a *fixed* separator character, then we get the right answer:

```
> strsplit( x = monkey, split = ".", fixed = TRUE )
[[1]]
[1] "It was the best of times"    " It was the blurst of times"
```

However, if we don't do this, then R will assume that when you typed `split = ". "` you were trying to construct a "regular expression", and as it happens the character `.` has a special meaning within a regular expression. As a consequence, if you forget to include the `fixed = TRUE` part, you won't get the answers you're looking for.

7.8.4 Making simple conversions

A slightly different task that comes up quite often is making transformations to text. A simple example of this would be converting text to lower case or upper case, which you can do using the `toupper()` and `tolower()` functions. Both of these functions have a single argument `x` which contains the text that needs to be converted. An example of this is shown below:

```
> text <- c( "lIfe", "Impact" )
> tolower( x = text )
[1] "life"    "impact"
```

A slightly more powerful way of doing text transformations is to use the `chartr()` function, which allows you to specify a "character by character" substitution. This function contains three arguments, `old`, `new` and `x`. As usual `x` specifies the text that needs to be transformed. The `old` and `new` arguments are strings of the same length, and they specify how `x` is to be converted. Every instance of the first character in `old` is converted to the first character in `new` and so on. For instance, suppose I wanted to convert `"albino"` to `"libido"`. To do this, I need to convert all of the `"a"` characters (all 1 of them) in `"albino"` into `"l"` characters (i.e., `a → l`). Additionally, I need to make the substitutions `l → i` and `n → d`. To do so, I would use the following command:

```
> old.text <- "albino"
> chartr( old = "aln", new = "lid", x = old.text )
[1] "libido"
```

7.8.5 Applying logical operations to text

In Section 3.8.5 we discussed a very basic text processing tool, namely the ability to use the equality operator `==` to test to see if two strings are identical to each other. However, you can also use other logical operators too. For instance R also allows you to use the `<` and `>` operators to determine which of two strings comes first, alphabetically speaking. Sort of. Actually, it's a bit more complicated than that, but let's start with a simple example:

```
> "cat" < "dog"
[1] TRUE
```

In this case, we see that `"cat"` does come before `"dog"` alphabetically, so R judges the statement to be true. However, if we ask R to tell us if `"cat"` comes before `"anteater"`,

```
> "cat" < "anteater"
[1] FALSE
```

It tells us that the statement is false. So far, so good. But text data is a bit more complicated than the dictionary suggests. What about `"cat"` and `"CAT"`? Which of these comes first? Let's try it and find out:

```
> "CAT" < "cat"
[1] TRUE
```

Table 7.3: The ordering of various text characters used by the `<` and `>` operators, as well as by the `sort()` function. Not shown is the “space” character, which actually comes first on the list.

```
! " # $ % & ' ( ) * + , - . / 0 1 2 3 4 5 6 7 8 9 : ; < = > ? @
A B C D E F G H I J K L M N O P Q R S T U V W X Y Z [ \ ] ^ _ '
a b c d e f g h i j k l m n o p q r s t u v w x y z } | {
```

.....

In other words, R assumes that uppercase letters come before lowercase ones. Fair enough. No-one is likely to be surprised by that. What you might find surprising is that R assumes that *all* uppercase letters come before *all* lowercase ones. That is, while "`anteater`" `<` "`zebra`" is a true statement, and the uppercase equivalent "`ANTEATER`" `<` "`ZEBRA`" is also true, it is *not* true to say that "`anteater`" `<` "`ZEBRA`", as the following extract illustrates:

```
> "anteater" < "ZEBRA"
[1] FALSE
```

This may seem slightly counterintuitive. With that in mind, it may help to have a quick look Table 7.3, which lists various text characters in the order that R uses.

7.8.6 Concatenating and printing with `cat()`

One function that I want to make a point of talking about, even though it’s not quite on topic, is the `cat()` function. The `cat()` function is a mixture of `paste()` and `print()`. That is, what it does is concatenate strings and then print them out. In your own work you can probably survive without it, since `print()` and `paste()` will actually do what you need, but the `cat()` function is so widely used that I think it’s a good idea to talk about it here. The basic idea behind `cat()` is straightforward. Like `paste()`, it takes several arguments as inputs, which it converts to strings, collapses (using a separator character specified using the `sep` argument), and prints on screen. If you want, you can use the `file` argument to tell R to print the output into a file rather than on screen (I won’t do that here). However, it’s important to note that the `cat()` function collapses vectors first, and *then* concatenates them. That is, notice that when I use `cat()` to combine `hw` and `ng`, I get a different result than if I’d used `paste()`

```
> cat( hw, ng )
hello world nasty government
> paste( hw, ng, collapse = " " )
[1] "hello nasty world government"
```

Notice the difference in the ordering of words. There’s a few additional details that I need to mention about `cat()`. Firstly, `cat()` really is a function for *printing*, and not for creating text strings to store for later. You can’t assign the output to a variable, as the following example illustrates:

```
> x <- cat( hw, ng )
hello world nasty government
> x
NULL
```

Despite my attempt to store the output as a variable, `cat()` printed the results on screen anyway, and it turns out that the variable I created doesn’t contain anything at all.¹⁹ Secondly, the `cat()` function

¹⁹Note that you can capture the output from `cat()` if you want to, but you have to be sneaky and use the `capture.output()` function. For example, the command `x <- capture.output(cat(hw,ng))` would work just fine.

makes use of a number of “special” characters. I’ll talk more about these in the next section, but I’ll illustrate the basic point now, using the example of “`\n`” which is interpreted as a “new line” character. For instance, compare the behaviour of `print()` and `cat()` when asked to print the string `hello\nworld`:

```
> print( "hello\nworld" ) # print literally:  
[1] "hello\nworld"  
> cat( "hello\nworld" ) # interpret as newline  
hello  
world
```

In fact, this behaviour is important enough that it deserves a section of its very own...

7.8.7 Using escape characters in text

The previous section brings us quite naturally to a fairly fundamental issue when dealing with strings, namely the issue of delimiters and escape characters. Reduced to its most basic form, the problem we have is that R commands are written using text characters, and our strings also consist of text characters. So, suppose I want to type in the word “hello”, and have R encode it as a string. If I were to just type `hello`, R will think that I’m referring to a variable or a function called `hello` rather than interpret it as a string. The solution that R adopts is to require you to enclose your string by **delimiter** characters, which can be either double quotes or single quotes. So, when I type `"hello"` or `'hello'` then R knows that it should treat the text in between the quote marks as a character string. However, this isn’t a complete solution to the problem: after all, “`”` and “`’`” are themselves perfectly legitimate text characters, and so we might want to include those in our string as well. For instance, suppose I wanted to encode the name “O’Rourke” as a string. It’s *not* legitimate for me to type `'O'rourke'` because R is too stupid to realise that “O’Rourke” is a real word. So it will interpret the `'O'` part as a complete string, and then will get confused when it reaches the `Rourke`’ part. As a consequence, what you get is an error message:

```
> 'O'Rourke'  
Error: unexpected symbol in "'O'Rourke"
```

To some extent, R offers us a cheap fix to the problem because of the fact that it allows us to use either “`”` or “`’`” as the delimiter character. Although `'O'rourke'` will make R cry, it is perfectly happy with `"O'Rourke"`:

```
> "O'Rourke"  
[1] "O'Rourke"
```

This is a real advantage to having two different delimiter characters. Unfortunately, anyone with even the slightest bit of deviousness to them can see the problem with this. Suppose I’m reading a book that contains the following passage,

P.J. O’Rourke says, “Yay, money!”. It’s a joke, but no-one laughs.

and I want to enter this as a string. Neither the `’` or `”` delimiters will solve the problem here, since this string contains both a single quote character and a double quote character. To encode strings like this one, we have to do something a little bit clever.

The solution to the problem is to designate an **escape character**, which in this case is `\`, the humble backslash. The escape character is a bit of a sacrificial lamb: if you include a backslash character in your string, R will *not* treat it as a literal character at all. It’s actually used as a way of inserting “special” characters into your string. For instance, if you want to force R to insert actual quote marks into the string, then what you actually type is `\'` or `\"` (these are called **escape sequences**). So, in order to encode the string discussed earlier, here’s a command I could use:

Table 7.4: Standard escape characters that are evaluated by some text processing commands, including `cat()`. This convention dates back to the development of the C programming language in the 1970s, and as a consequence a lot of these characters make most sense if you pretend that R is actually a typewriter, as explained in the main text. Type `?Quotes` for the corresponding R help file.

Escape sequence	Interpretation	Escape sequence	Interpretation
\n	Newline	\f	Form feed
\t	Horizontal Tab	\a	Alert sound
\v	Vertical Tab	\\\	Backslash
\b	Backspace	\'	Single quote
\r	Carriage Return	\"	Double quote

```
> PJ <- "P.J. O'Rourke says, \"Yay, money!\". It's a joke, but no-one laughs."
```

Notice that I've included the backslashes for both the single quotes and double quotes. That's actually overkill: since I've used `"` as my delimiter, I only needed to do this for the double quotes. Nevertheless, the command has worked, since I didn't get an error message. Now let's see what happens when I print it out:

```
> print( PJ )
[1] "P.J. O'Rourke says, \"Yay, money!\". It's a joke, but no-one laughs."
```

Hm. Why has R printed out the string using `\"`? For the exact same reason that *I* needed to insert the backslash in the first place. That is, when R prints out the `PJ` string, it has enclosed it with delimiter characters, and it wants to unambiguously show us which of the double quotes are delimiters and which ones are actually part of the string. Fortunately, if this bugs you, you can make it go away by using the `print.noquote()` function, which will just print out the literal string that you encoded in the first place:

```
> print.noquote( PJ )
[1] P.J. O'Rourke says, "Yay, money!". It's a joke, but no-one laughs.
```

Typing `cat(PJ)` will produce a similar output.

Introducing the escape character solves a lot of problems, since it provides a mechanism by which we can insert all sorts of characters that aren't on the keyboard. For instance, as far as a computer is concerned, “new line” is actually a text character. It's the character that is printed whenever you hit the “return” key on your keyboard. If you want to insert a new line character into your string, you can actually do this by including the escape sequence `\n`. Or, if you want to insert a backslash character, then you can use `\\\`. A list of the standard escape sequences recognised by R is shown in Table 7.4. A lot of these actually date back to the days of the typewriter (e.g., carriage return), so they might seem a bit counterintuitive to people who've never used one. In order to get a sense for what the various escape sequences do, we'll have to use the `cat()` function, because it's the only function “dumb” enough to literally print them out:

```
> cat( "xxxx\boo" ) # \b is a backspace, so it deletes the preceding x
xxxx
> cat( "xxxx\t oo" ) # \t is a tab, so it inserts a tab space
xxxx      oo
> cat( "xxxx\n oo" ) # \n is a newline character
xxxx
oo
```

```
> cat( "xxxx\roo" ) # \r returns you to the beginning of the line
oooo
```

And that's pretty much it. There are a few other escape sequence that R recognises, which you can use to insert arbitrary ASCII or Unicode characters into your string (type `?Quotes` for more details) but I won't go into details here.

7.8.8 Matching and substituting text

Another task that we often want to solve is find all strings that match a certain criterion, and possibly even to make alterations to the text on that basis. There are several functions in R that allow you to do this, three of which I'll talk about briefly here: `grep()`, `gsub()` and `sub()`. Much like the `substr()` function that I talked about earlier, all three of these functions are intended to be used in conjunction with regular expressions (see Section 7.8.9) but you can also use them in a simpler fashion, since they all allow you to set `fixed = TRUE`, which means we can ignore all this regular expression rubbish and just use simple text matching.

So, how do these functions work? Let's start with the `grep()` function. The purpose of this function is to input a vector of character strings `x`, and to extract all those strings that fit a certain pattern. In our examples, I'll assume that the `pattern` in question is a literal sequence of characters that the string must contain (that's what `fixed = TRUE` does). To illustrate this, let's start with a simple data set, a vector that contains the names of three `beers`. Something like this:

```
> beers <- c( "little creatures", "sierra nevada", "coopers pale" )
```

Next, let's use `grep()` to find out which of these strings contains the substring "`er`". That is, the `pattern` that we need to match is the fixed string "`er`", so the command we need to use is:

```
> grep( pattern = "er", x = beers, fixed = TRUE )
[1] 2 3
```

What the output here is telling us is that the second and third elements of `beers` both contain the substring "`er`". Alternatively, however, we might prefer it if `grep()` returned the actual strings themselves. We can do this by specifying `value = TRUE` in our function call. That is, we'd use a command like this:

```
> grep( pattern = "er", x = beers, fixed = TRUE, value = TRUE )
[1] "sierra nevada" "coopers pale"
```

The other two functions that I wanted to mention in this section are `gsub()` and `sub()`. These are both similar in spirit to `grep()` insofar as what they do is search through the input strings (`x`) and find all of the strings that match a `pattern`. However, what these two functions do is *replace* the pattern with a `replacement` string. The `gsub()` function will replace *all* instances of the pattern, whereas the `sub()` function just replaces the first instance of it in each string. To illustrate how this works, suppose I want to replace all instances of the letter "`a`" with the string "`BLAH`". I can do this to the `beers` data using the `gsub()` function:

```
> gsub( pattern = "a", replacement = "BLAH", x = beers, fixed = TRUE )
[1] "little creBLAHtures"      "sierrBLAH nevBLAHdBLAH"
[3] "coopers pBLAHle"
```

Notice that all three of the "`a`'s in "`sierra nevada`" have been replaced. In contrast, let's see what happens when we use the exact same command, but this time using the `sub()` function instead:

```
> sub( pattern = "a", replacement = "BLAH", x = beers, fixed = TRUE )
[1] "little creBLAHtures" "sierrBLAH nevada"      "coopers pBLAHle"
```

Only the first "a" is changed.

7.8.9 Regular expressions (not really)

There's one last thing I want to talk about regarding text manipulation, and that's the concept of a **regular expression**. Throughout this section we've often needed to specify `fixed = TRUE` in order to force R to treat some of our strings as actual strings, rather than as regular expressions. So, before moving on, I want to very briefly explain what regular expressions are. I'm *not* going to talk at all about how they work or how you specify them, because they're genuinely complicated and not at all relevant to this book. However, they are extremely powerful tools and they're quite widely used by people who have to work with lots of text data (e.g., people who work with natural language data), and so it's handy to at least have a vague idea about what they are. The basic idea is quite simple. Suppose I want to extract all strings in my `beers` vector that contain a vowel followed immediately by the letter "s". That is, I want to find the beer names that contain either "as", "es", "is", "os" or "us". One possibility would be to manually specify all of these possibilities and then match against these as fixed strings one at a time, but that's tedious. The alternative is to try to write out a single "regular" expression that matches all of these. The regular expression that does this²⁰ is "[aeiou]s", and you can kind of see what the syntax is doing here. The bracketed expression means "any of the things in the middle", so the expression as a whole means "any of the things in the middle" (i.e. vowels) followed by the letter "s". When applied to our beer names we get this:

```
> grep( pattern = "[aeiou]s", x = beers, value = TRUE )
[1] "little creatures"
```

So it turns out that only "little creatures" contains a vowel followed by the letter "s". But of course, had the data contained a beer like "fosters", that would have matched as well because it contains the string "os". However, I deliberately chose not to include it because Fosters is not – in my opinion – a proper beer.²¹ As you can tell from this example, regular expressions are a neat tool for specifying *patterns* in text: in this case, "vowel then s". So they are definitely things worth knowing about if you ever find yourself needing to work with a large body of text. However, since they are fairly complex and not necessary for any of the applications discussed in this book, I won't talk about them any further.

7.9

Reading unusual data files

In this section I'm going to switch topics (again!) and turn to the question of how you can load data from a range of different sources. Throughout this book I've assumed that your data are stored as an `.Rdata` file or as a "properly" formatted CSV file. And if so, then the basic tools that I discussed in Section 4.5 should be quite sufficient. However, in real life that's not a terribly plausible assumption to make, so I'd better talk about some of the other possibilities that you might run into.

²⁰Sigh. For advanced users: R actually supports two different ways of specifying regular expressions. One is the POSIX standard, the other is to use Perl-style regular expressions. The default is generally POSIX. If you understand regular expressions, that probably made sense to you. If not, don't worry. It's not important.

²¹I thank Amy Perfors for this example.

7.9.1 Loading data from text files

The first thing I should point out is that if your data are saved as a text file but aren't *quite* in the proper CSV format, then there's still a pretty good chance that the `read.csv()` function (or equivalently, `read.table()`) will be able to open it. You just need to specify a few more of the optional arguments to the function. If you type `?read.csv` you'll see that the `read.csv()` function actually has several arguments that you can specify. Obviously you need to specify the `file` that you want it to load, but the others all have sensible default values. Nevertheless, you will sometimes need to change them. The ones that I've often found myself needing to change are:

- `header`. A lot of the time when you're storing data as a CSV file, the first row actually contains the column names and not data. If that's not true, you need to set `header = FALSE`.
- `sep`. As the name "comma separated value" indicates, the values in a row of a CSV file are usually separated by commas. This isn't universal, however. In Europe the decimal point is typically written as `,` instead of `.` and as a consequence it would be somewhat awkward to use `,` as the separator. Therefore it is not unusual to use `;` over there. At other times, I've seen a TAB character used. To handle these cases, we'd need to set `sep = ";"` or `sep = "\t"`.
- `quote`. It's conventional in CSV files to include a quoting character for textual data. As you can see by looking at the `booksales.csv` file, this is usually a double quote character, `"`. But sometimes there is no quoting character at all, or you might see a single quote mark `'` used instead. In those cases you'd need to specify `quote = ""` or `quote = "'"`.
- `skip`. It's actually very common to receive CSV files in which the first few rows have nothing to do with the actual data. Instead, they provide a human readable summary of where the data came from, or maybe they include some technical info that doesn't relate to the data. To tell R to ignore the first (say) three lines, you'd need to set `skip = 3`.
- `na.strings`. Often you'll get given data with missing values. For one reason or another, some entries in the table are missing. The data file needs to include a "special" string to indicate that the entry is missing. By default R assumes that this string is `NA`, since that's what *it* would do, but there's no universal agreement on what to use in this situation. If the file uses `???` instead, then you'll need to set `na.strings = "???"`.

It's kind of nice to be able to have all these options that you can tinker with. For instance, have a look at the data file shown pictured in Figure 7.1. This file contains almost the same data as the last file (except it doesn't have a header), and it uses a bunch of wacky features that you don't normally see in CSV files. In fact, it just so happens that I'm going to have to change all five of those arguments listed above in order to load this file. Here's how I would do it:

```
> data <- read.csv( file = "booksales2.csv", # specify the name of the file
+                     header = FALSE,          # variable names in the file?
+                     skip = 8,                # ignore the first 8 lines
+                     quote = "*",             # what indicates text data?
+                     sep = "\t",              # what separates different entries?
+                     na.strings = "NFI" )     # what is the code for missing data?
```

If I now have a look at the data I've loaded, I see that this is what I've got:

```
> head( data )
      V1  V2  V3   V4
1  January 31    0 high
```

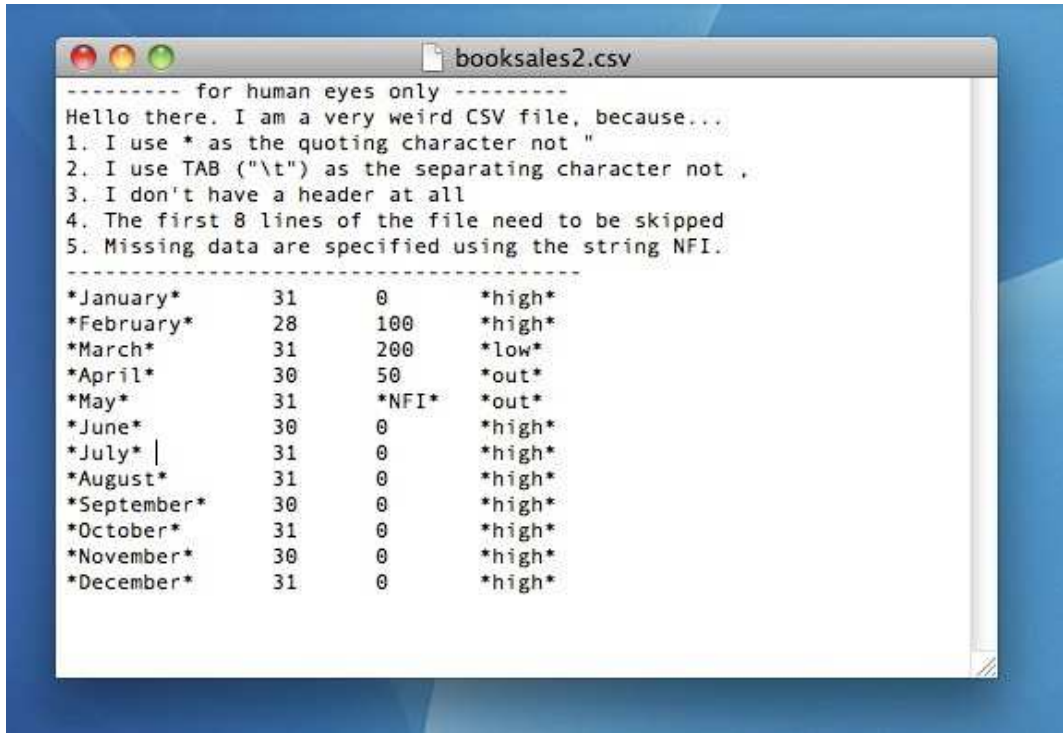


Figure 7.1: The `booksales2.csv` data file. It contains more or less the same data as the original `booksales.csv` data file, but has a lot of very quirky features.

```

2 February 28 100 high
3 March 31 200 low
4 April 30 50 out
5 May 31 NA out
6 June 30 0 high

```

Because I told R to expect `*` to be used as the quoting character instead of `"`; to look for tabs (which we write like this: `\t`) instead of commas, and to skip the first 8 lines of the file, it's basically loaded the right data. However, since `booksales2.csv` doesn't contain the column names, R has made them up. Showing the kind of imagination I expect from insentient software, R decided to call them `V1`, `V2`, `V3` and `V4`. Finally, because I told it that the file uses "NFI" to denote missing data, R correctly figures out that the sales data for May are actually missing.

In real life you'll rarely see data this stupidly formatted.²²

7.9.2 Loading data from SPSS (and other statistics packages)

The commands listed above are the main ones we'll need for data files in this book. But in real life we have many more possibilities. For example, you might want to read data files in from other statistics programs. Since SPSS is probably the most widely used statistics package in psychology, it's worth

²²If you're lucky.

briefly showing how to open SPSS data files (file extension `.sav`). It's surprisingly easy. The extract below should illustrate how to do so:

```
> library( foreign )          # load the package
> X <- read.spss( "datafile.sav" )  # create a list containing the data
> X <- as.data.frame( X )        # convert to data frame
```

If you wanted to import from an SPSS file to a data frame directly, instead of importing a list and then converting the list to a data frame, you can do that too:

```
> X <- read.spss( file = "datafile.sav", to.data.frame = TRUE )
```

And that's pretty much it, at least as far as SPSS goes. As far as other statistical software goes, the `foreign` package provides a wealth of possibilities. To open SAS files, check out the `read.ssd()` and `read.xport()` functions. To open data from Minitab, the `read.mtp()` function is what you're looking for. For Stata, the `read.dta()` function is what you want. For Systat, the `read.systat()` function is what you're after.

7.9.3 Loading Excel files

A different problem is posed by Excel files. Despite years of yelling at people for sending data to me encoded in a proprietary data format, I get sent a lot of Excel files. In general R does a pretty good job of opening them, but it's bit finicky because Microsoft don't seem to be terribly fond of people using non-Microsoft products, and go to some lengths to make it tricky. If you get an Excel file, my suggestion would be to open it up in Excel (or better yet, OpenOffice, since that's free software) and then save the spreadsheet as a CSV file. Once you've got the data in that format, you can open it using `read.csv()`. However, if for some reason you're desperate to open the `.xls` or `.xlsx` file directly, then you can use the `read.xls()` function in the `gdata` package:

```
> library( gdata )           # load the package
> X <- read.xls( "datafile.xlsx" )  # create a data frame
```

This usually works. And if it doesn't, you're probably justified in "suggesting" to the person that sent you the file that they should send you a nice clean CSV file instead.

7.9.4 Loading Matlab (& Octave) files

A lot of scientific labs use Matlab as their default platform for scientific computing; or Octave as a free alternative. Opening Matlab data files (file extension `.mat`) slightly more complicated, and if it wasn't for the fact that Matlab is so very widespread and is an extremely good platform, I wouldn't mention it. However, since Matlab is so widely used, I think it's worth discussing briefly how to get Matlab and R to play nicely together. The way to do this is to install the `R.matlab` package (don't forget to install the dependencies too). Once you've installed and loaded the package, you have access to the `readMat()` function. As any Matlab user will know, the `.mat` files that Matlab produces are workspace files, very much like the `.Rdata` files that R produces. So you can't import a `.mat` file as a data frame. However, you can import it as a list. So, when we do this:

```
> library( R.matlab )          # load the package
> data <- readMat( "matlabfile.mat" )  # read the data file to a list
```

The `data` object that gets created will be a list, containing one variable for every variable stored in the Matlab file. It's fairly straightforward, though there are some subtleties that I'm ignoring. In particular,

note that if you don't have the `Rcompression` package, you can't open Matlab files above the version 6 format. So, if like me you've got a recent version of Matlab, and don't have the `Rcompression` package, you'll need to save your files using the `-v6` flag otherwise R can't open them.

Oh, and Octave users? The `foreign` package contains a `read.octave()` command. Just this once, the world makes life easier for you folks than it does for all those cashed-up swanky Matlab bastards.

7.9.5 Saving other kinds of data

Given that I talked extensively about how to load data from non-R files, it might be worth briefly mentioning that R is also pretty good at writing data into other file formats besides its own native ones. I won't discuss them in this book, but the `write.csv()` function can write CSV files, and the `write.foreign()` function (in the `foreign` package) can write SPSS, Stata and SAS files. There are also a lot of low level commands that you can use to write very specific information to a file, so if you really, really needed to you could create your own `write.obscurefiletype()` function, but that's also a long way beyond the scope of this book. For now, all that I want you to recognise is that this capability is there if you need it.

7.9.6 Are we done yet?

Of course not. If I've learned nothing else about R it's that you're *never bloody done*. This listing doesn't even come close to exhausting the possibilities. Databases are supported by the `RODBC`, `DBI`, and `RMySQL` packages among others. You can open webpages using the `RCurl` package. Reading and writing JSON objects is supported through the `rjson` package. And so on. In a sense, the right question is not so much "can R do this?" so much as "whereabouts in the wilds of CRAN *is* the damn package that does it?"

7.10

Coercing data from one class to another

Sometimes you want to change the variable class. This can happen for all sorts of reasons. Sometimes when you import data from files, it can come to you in the wrong format: numbers sometimes get imported as text, dates usually get imported as text, and many other possibilities besides. Regardless of how you've ended up in this situation, there's a very good chance that sometimes you'll want to convert a variable from one class into another one. Or, to use the correct term, you want to `coerce` the variable from one class into another. Coercion is a little tricky, and so I'll only discuss the very basics here, using a few simple examples.

Firstly, let's suppose we have a variable `x` that is *supposed* to be representing a number, but the data file that you've been given has encoded it as text. Let's imagine that the variable is something like this:

```
> x <- "100" # the variable
> class(x)    # what class is it?
[1] "character"
```

Obviously, if I want to do calculations using `x` in its current state, R is going to get very annoyed at me. It thinks that `x` is text, so it's not going to allow me to try to do mathematics using it! Obviously, we need to coerce `x` from character to numeric. We can do that in a straightforward way by using the `as.numeric()` function:

```

> x <- as.numeric(x) # coerce the variable
> class(x)           # what class is it?
[1] "numeric"
> x + 1              # hey, addition works!
[1] 101

```

Not surprisingly, we can also convert it back again if we need to. The function that we use to do this is the `as.character()` function:

```

> x <- as.character(x) # coerce back to text
> class(x)            # check the class:
[1] "character"

```

However, there's some fairly obvious limitations: you can't coerce the string "`hello world`" into a number because, well, there's isn't a number that corresponds to it. Or, at least, you can't do anything useful:

```

> as.numeric( "hello world" ) # this isn't going to work.
[1] NA
Warning message:
NAs introduced by coercion

```

In this case R doesn't give you an error message; it just gives you a warning, and then says that the data is missing (see Section 4.6.1 for the interpretation of `NA`).

That gives you a feel for how to change between numeric and character data. What about logical data? To cover this briefly, coercing text to logical data is pretty intuitive: you use the `as.logical()` function, and the character strings "`T`", "`TRUE`", "`True`" and "`true`" all convert to the logical value of `TRUE`. Similarly "`F`", "`FALSE`", "`False`", and "`false`" all become `FALSE`. All other strings convert to `NA`. When you go back the other way using `as.character()`, `TRUE` converts to "`TRUE`" and `FALSE` converts to "`FALSE`". Converting numbers to logicals – again using `as.logical()` – is straightforward. Following the convention in the study of logic, the number `0` converts to `FALSE`. Everything else is `TRUE`. Going back using `as.numeric()`, `FALSE` converts to `0` and `TRUE` converts to `1`.

7.11

Other useful data structures

Up to this point we have encountered several different kinds of variables. At the simplest level, we've seen numeric data, logical data and character data. However, we've also encountered some more complicated kinds of variables, namely factors, formulas, data frames and lists. We'll see a few more specialised data structures later on in this book, but there's a few more generic ones that I want to talk about in passing. None of them are central to the rest of the book (and in fact, the only one we'll even see anywhere else is the matrix), but they do crop up a fair bit in real life.

7.11.1 Matrices (and arrays)

In various different places in this chapter I've made reference to an R data structure called a `matrix`, and mentioned that I'd talk a bit more about matrices later on. That time has come. Much like a data frame, a matrix is basically a big rectangular table of data, and in fact there are quite a few similarities

between the two. However, there are also some key differences, so it's important to talk about matrices in a little detail. Let's start by using `rbind()` to create a small matrix:²³

```
> row.1 <- c( 2,3,1 )           # create data for row 1
> row.2 <- c( 5,6,7 )           # create data for row 2
> M <- rbind( row.1, row.2 )    # row bind them into a matrix
> print( M )                   # and print it out...
[,1] [,2] [,3]
row.1     2     3     1
row.2     5     6     7
```

The variable `M` is a matrix, which we can confirm by using the `class()` function. Notice that, when we bound the two vectors together, R retained the names of the original variables as row names. We could delete these if we wanted by typing `rownames(M)<-NULL`, but I generally prefer having meaningful names attached to my variables, so I'll keep them. In fact, let's also add some highly unimaginative column names as well:

```
> colnames(M) <- c( "col.1", "col.2", "col.3" )
> print(M)
  col.1 col.2 col.3
row.1     2     3     1
row.2     5     6     7
```

You can use square brackets to subset a matrix in much the same way that you can for data frames, again specifying a row index and then a column index. For instance, `M[2,3]` pulls out the entry in the 2nd row and 3rd column of the matrix (i.e., 7), whereas `M[2,]` pulls out the entire 2nd row, and `M[,3]` pulls out the entire 3rd column. However, it's worth noting that when you pull out a column, R will print the results horizontally, not vertically. The reason for this relates to how matrices (and arrays generally) are implemented. The original matrix `M` is treated as a two-dimensional objects, containing 2 rows and 3 columns. However, whenever you pull out a single row or a single column, the result is considered to be one-dimensional. As far as R is concerned there's no real reason to distinguish between a one-dimensional object printed vertically (a column) and a one-dimensional object printed horizontally (a row), and it prints them all out horizontally.²⁴ There is also a way of using only a single index, but due to the internal structure to how R defines a matrix, it works very differently to what we saw previously with data frames. The single-index approach is illustrated in Table 7.5 but I don't really want to focus on it since we'll never really need it for this book, and matrices don't play anywhere near as large a role in this book as data frames do.

The critical difference between a data frame and a matrix is that, in a data frame, we have this notion that each of the columns corresponds to a different variable: as a consequence, the columns in a data frame can be of different data types. The first column could be numeric, and the second column could contain character strings, and the third column could be logical data. In that sense, there is a fundamental asymmetry build into a data frame, because of the fact that columns represent variables

²³You can also use the `matrix()` command itself, but I think the “binding” approach is a little more intuitive.

²⁴This has some interesting implications for how matrix algebra is implemented in R (which I'll admit I initially found odd), but that's a little beyond the scope of this book. However, since there will be a small proportion of readers that do care, I'll quickly outline the basic thing you need to get used to: when multiplying a matrix by a vector (or one-dimensional array) using the `%^%` operator R will attempt to interpret the vector (or 1D array) as either a row-vector or column-vector, depending on whichever one makes the multiplication work. That is, suppose `M` is the 2×3 matrix, and `v` is a 1×3 row vector. It is impossible to multiply `Mv`, since the dimensions don't conform, but you *can* multiply by the corresponding column vector, `Mv'`. So, if I set `v <- M[2,]` and then try to calculate `M %^% v`, which you'd think would fail, it actually works because R treats the one dimensional array as if it were a column vector for the purposes of matrix multiplication. Note that if both objects are one dimensional arrays/vectors, this leads to ambiguity since vv' (inner product) and $v'v$ (outer product) yield different answers. In this situation, the `%^%` operator returns the inner product not the outer product. To understand all the details, check out the help documentation.

Table 7.5: An illustration of two different ways of indexing a 2×3 matrix. On the left we see the row and column version, which is identical to the corresponding indexing scheme for a data frame of the same size. On the right we see the single-index version, which is quite different to what we would get with a data frame. The reason for this is that, for both data frames and matrices, the “row and column” version exists to allow the human user to interact with the object in the psychologically meaningful way: since both data frames and matrices are basically just tables of data, it’s the same in each case. However, the single-index version is really a method for you to interact with the object in terms of its internal structure, and the internals for data frames and matrices are quite different.

row	column			row	column		
	1	2	3		1	2	3
1	[1,1]	[1,2]	[1,3]	1	[1]	[3]	[5]
2	[2,1]	[2,2]	[2,3]	2	[2]	[4]	[6]
.....							

(which can be qualitatively different to each other) and rows represent cases (which cannot). Matrices are intended to be thought of in a different way. At a fundamental level, a matrix really is just one variable: it just happens that this one variable is formatted into rows and columns. If you want a matrix of numeric data, every single element in the matrix *must* be a number. If you want a matrix of character strings, every single element in the matrix *must* be a character string. If you try to mix data of different types together, then R will either spit out an error, or quietly coerce the underlying data into a list. If you want to find out what class R secretly thinks the data within the matrix is, you need to do something like this:

```
> class( M[1] )
[1] "numeric"
```

You can’t type `class(M)`, because all that will happen is R will tell you that `M` is a matrix: we’re not interested in the class of the matrix itself, we want to know what class the underlying data is assumed to be. Anyway, to give you a sense of how R enforces this, let’s try to change one of the elements of our numeric matrix into a character string:

```
> M[1,2] <- "text"
> M
   col.1 col.2 col.3
row.1 "2"   "text" "1"
row.2 "5"   "6"   "7"
```

It looks as if R has coerced all of the data in our matrix into character strings. And in fact, if we now typed in `class(M[1])` we’d see that this is exactly what has happened. If you alter the contents of one element in a matrix, R will change the underlying data type as necessary.

There’s only one more thing I want to talk about regarding matrices. The concept behind a matrix is very much a mathematical one, and in mathematics a matrix is a most definitely a two-dimensional object. However, when doing data analysis, we often have reasons to want to use higher dimensional tables (e.g., sometimes you need to cross-tabulate three variables against each other). You can’t do this with matrices, but you can do it with `arrays`. An array is just like a matrix, except it can have more than two dimensions if you need it to. In fact, as far as R is concerned a matrix is just a special kind of array, in much the same way that a data frame is a special kind of list. I don’t want to talk about arrays too much, but I will very briefly show you an example of what a 3D array looks like. To that end, let’s cross tabulate the `speaker` and `utterance` variables from the `nightgarden.Rdata` data file, but we’ll add

a third variable to the cross-tabs this time, a logical variable which indicates whether or not I was still awake at this point in the show:

```
> dan.awake <- c( TRUE, TRUE, TRUE, TRUE, TRUE, FALSE, FALSE, FALSE, FALSE )
```

Now that we've got all three variables in the workspace (assuming you loaded the `nightgarden.Rdata` data earlier in the chapter) we can construct our three way cross-tabulation, using the `table()` function.

```
> xtab.3d <- table( speaker, utterance, dan.awake )
> xtab.3d
, , dan.awake = FALSE

      utterance
speaker    ee onk oo pip
makka-pakka 0   2   0   2
tombliboo   0   0   1   0
upsy-daisy  0   0   0   0

, , dan.awake = TRUE

      utterance
speaker    ee onk oo pip
makka-pakka 0   0   0   0
tombliboo   1   0   0   0
upsy-daisy  0   2   0   2
```

Hopefully this output is fairly straightforward: because R can't print out text in three dimensions, what it does is show a sequence of 2D slices through the 3D table. That is, the `, , dan.awake = FALSE` part indicates that the 2D table that follows below shows the 2D cross-tabulation of `speaker` against `utterance` only for the `dan.awake = FALSE` instances, and so on.²⁵

7.11.2 Ordered factors

One topic that I neglected to mention when discussing factors previously (Section 4.7) is that there are actually two different types of factor in R, unordered factors and ordered factors. An unordered factor corresponds to a nominal scale variable, and all of the factors we've discussed so far in this book have been unordered (as will all the factors used anywhere else except in this section). However, it's often very useful to explicitly tell R that your variable is *ordinal scale*, and if so you need to declare it to be an **ordered factor**. For instance, earlier in this chapter we made use of a variable consisting of Likert scale data, which we represented as the `likert.raw` variable:

```
> likert.raw
[1] 1 7 3 4 4 4 2 6 5 5
```

We can declare this to be an ordered factor in by using the `factor()` function, and setting `ordered = TRUE`. To illustrate how this works, let's create an ordered factor called `likert.ordinal` and have a look at it:

²⁵I should note that if you type `class(xtab.3d)` you'll discover that this is a "table" object rather than an "array" object. However, this labelling is only skin deep. The underlying data structure here is actually an array. Advanced users may wish to check this using the command `class(unclass(xtab.3d))`, but it's not important for our purposes. All I really want to do in this section is show you what the output looks like when you encounter a 3D array.

```

> likert.ordinal <- factor( x = likert.raw,           # the raw data
+                           levels = seq(7,1,-1), # strongest agreement is 1, weakest is 7
+                           ordered = TRUE )      # and it's ordered
> print( likert.ordinal )
[1] 1 7 3 4 4 4 2 6 5 5
Levels: 7 < 6 < 5 < 4 < 3 < 2 < 1

```

Notice that when we print out the ordered factor, R explicitly tells us what order the levels come in. Because I wanted to order my levels in terms of *increasing* strength of agreement, and because a response of 1 corresponded to the strongest agreement and 7 to the strongest disagreement, it was important that I tell R to encode 7 as the lowest value and 1 as the largest. Always check this when creating an ordered factor: it's very easy to accidentally encode your data "upside down" if you're not paying attention. In any case, note that we can (and should) attach meaningful names to these factor levels by using the `levels()` function, like this:

```

> levels( likert.ordinal ) <- c( "strong.disagree", "disagree", "weak.disagree",
+                               "neutral", "weak.agree", "agree", "strong.agree" )
> print( likert.ordinal )
[1] strong.agree    strong.disagree weak.agree       neutral        neutral        neutral
[7] agree          disagree        weak.disagree   weak.disagree
Levels: strong.disagree < disagree < weak.disagree < neutral < weak.agree < agree < strong.agree

```

One nice thing about using ordered factors is that there are a lot of analyses for which R automatically treats ordered factors differently from unordered factors, and generally in a way that is more appropriate for ordinal data. However, since I don't discuss that in this book, I won't go into details. Like so many things in this chapter, my main goal here is to make you aware that R has this capability built into it; so if you ever need to start thinking about interval scale variables in more detail, you have at least some idea where to start looking!

7.11.3 Dates and times

Times and dates are very annoying types of data. To a first approximation we can say that there are 365 days in a year, 24 hours in a day, 60 minutes in an hour and 60 seconds in a minute, but that's not quite correct. The length of the solar day is not exactly 24 hours, and the length of solar year is not exactly 365 days, so we have a complicated system of corrections that have to be made to keep the time and date system working. On top of that, the measurement of time is usually taken relative to a local time zone, and most (but not all) time zones have both a standard time and a daylight savings time, though the date at which the switch occurs is not at all standardised. So, as a form of data, times and dates *suck*. Unfortunately, they're also important. Sometimes it's possible to avoid having to use any complicated system for dealing with times and dates. Often you just want to know what year something happened in, so you can just use numeric data: in quite a lot of situations something as simple as `this.year <- 2011` works just fine. If you can get away with that for your application, this is probably the best thing to do. However, sometimes you really do need to know the actual date. Or, even worse, the actual time. In this section, I'll very briefly introduce you to the basics of how R deals with date and time data. As with a lot of things in this chapter, I won't go into details because I don't use this kind of data anywhere else in the book. The goal here is to show you the basics of what you need to do if you ever encounter this kind of data in real life. And then we'll all agree never to speak of it again.

To start with, let's talk about the date. As it happens, modern operating systems are very good at keeping track of the time and date, and can even handle all those annoying timezone issues and daylight savings pretty well. So R takes the quite sensible view that it can just ask the operating system what the date is. We can pull the date using the `Sys.Date()` function:

```
> today <- Sys.Date() # ask the operating system for the date
> print(today)          # display the date
[1] "2011-08-24"
```

Okay, that seems straightforward. But, it does rather look like `today` is just a character string, doesn't it? That would be a problem, because dates really do have a numeric character to them, and it would be nice to be able to do basic addition and subtraction to them. Well, fear not. If you type in `class(today)`, R will tell you that the class of the `today` variable is "`Date`". What this means is that, hidden underneath this text string that prints out an actual date, R actually has a numeric representation.²⁶ What that means is that you actually can add and subtract days. For instance, if we add 1 to `today`, R will print out the date for tomorrow:

```
> today + 1
[1] "2011-08-25"
```

Let's see what happens when we add 365 days:

```
> today + 365
[1] "2012-08-23"
```

It turns out that R handles this better than I would have, since I'd forgotten that 2012 is a leap year and so I would have gotten the wrong answer had I done it myself. R provides a number of functions for working with dates, but I don't want to talk about them in any detail. I will, however, make passing mention of the `weekdays()` function which will tell you what day of the week a particular date corresponded to, which is extremely convenient in some situations:

```
> weekdays( today )
[1] "Wednesday"
```

I'll also point out that you can use the `as.Date()` to convert various different kinds of data into dates. If the data happen to be strings formatted exactly according to the international standard notation (i.e., YYYY-MM-DD) then the conversion is straightforward, because that's the format that R expects to see by default. You can convert dates from other formats too, but it's slightly trickier, and beyond the scope of this book.

What about times? Well, times are even more annoying, so much so that I don't intend to talk about them at all in this book, other than to point you in the direction of some vaguely useful things. R itself does provide you with some tools for handling time data, and in fact there are two separate classes of data that are used to represent times, known by the odd names `POSIXct` and `POSIXlt`. You can use these to work with times if you want to, but for most applications you would probably be better off downloading the `chron` package, which provides some much more user friendly tools for working with times and dates.

7.12

Miscellaneous topics

To finish this chapter, I have a few topics to discuss that don't really fit in with any of the other things in this chapter. They're all kind of useful things to know about, but they are really just "odd topics" that don't fit with the other examples. Here goes:

²⁶Date objects are coded as the number of days that have passed since January 1, 1970.

7.12.1 The problems with floating point arithmetic

If I've learned nothing else about transfinite arithmetic (and I haven't) it's that infinity is a tedious and inconvenient concept. Not only is it annoying and counterintuitive at times, but it has nasty practical consequences. As we were all taught in high school, there are some numbers that *cannot* be represented as a decimal number of finite length, nor can they be represented as any kind of fraction between two whole numbers; $\sqrt{2}$, π and e , for instance. In everyday life we mostly don't care about this. I'm perfectly happy to approximate π as 3.14, quite frankly. Sure, this does produce some rounding errors from time to time, and if I'd used a more detailed approximation like 3.1415926535 I'd be less likely to run into those issues, but in all honesty I've never needed my calculations to be *that* precise. In other words, although our pencil and paper calculations cannot represent the number π exactly as a decimal number, we humans are smart enough to realise that we don't care. Computers, unfortunately, are dumb ... and you don't have to dig too deep in order to run into some very weird issues that arise because they can't represent numbers perfectly. Here is my favourite example:

```
> (0.1 * 3) == 0.3
[1] FALSE
```

Obviously, R has made a mistake here, because this is definitely the wrong answer. Your first thought might be that R is broken. But you can reproduce the same error in dozens of different programming languages, so the issue isn't specific to R. Your next thought might be that it's something in the hardware, but you can get the same mistake on any machine. It's something deeper than that.

The fundamental issue at hand is **floating point arithmetic**, which is a fancy way of saying that computers will *always* round a number to fixed number of significant digits. The exact number of significant digits that the computer stores isn't important to us:²⁷ what matters is that whenever the number that the computer is trying to store is very long, you get rounding errors. That's actually what's happening with our example above. There are teeny tiny rounding errors that have appeared in the computer's storage of the numbers, and these rounding errors have in turn caused the internal storage of `0.1 * 3` to be a tiny bit different from the internal storage of `0.3`. How big are these differences? Let's ask R:

```
> (0.1 * 3) - 0.3
[1] 5.551115e-17
```

Very tiny indeed. No sane person would care about differences that small. But R is not a sane person, and the equality operator `==` is very literal minded. It returns a value of `TRUE` only when the two values that it is given are absolutely identical to each other. And in this case they are not. However, this only answers half of the question. The other half of the question is, why are we getting these rounding errors when we're only using nice simple numbers like 0.1 and 0.3? This seems a little counterintuitive. The answer is that, like most programming languages, R doesn't store numbers using their *decimal* expansion (i.e., base 10: using digits 0, 1, 2 ..., 9). We humans like to write our numbers in base 10 because we have 10 fingers. But computers don't have fingers, they have transistors; and transistors are built to store 2 numbers not 10. So you can see where this is going: the internal storage of a number in R is based on its *binary* expansion (i.e., base 2: using digits 0 and 1). And unfortunately, here's what the binary expansion of 0.1 looks like:

$$.1(\text{decimal}) = .00011001100110011\dots(\text{binary})$$

²⁷For advanced users: type `?double` for more information.

and the pattern continues forever. In other words, from the perspective of your computer, which likes to encode numbers in binary,²⁸ the number 0.1 is not at all simple. It's actually an infinitely long binary number! As a consequence, the computer can make minor errors when doing calculations here.

With any luck you now understand the problem, which ultimately comes down to the twin fact that (1) we usually think in decimal numbers and computers usually compute with binary numbers, and (2) computers are finite machines and can't store infinitely long numbers. The only questions that remain are when you should care and what you should do about it. Thankfully, you don't have to care very often: because the rounding errors are small, the only practical situation that I've seen this issue arise for is when you want to test whether an arithmetic fact holds exactly (e.g., is someone's response time equal to *exactly* 2×0.33 seconds?) This is pretty rare in real world data analysis, but just in case it does occur, it's better to use a test that allows for a small *tolerance*. That is, if the difference between the two numbers is below a certain threshold value, we deem them to be equal for all practical purposes. For instance, you could do something like this, which asks whether the difference between the two numbers is less than a tolerance of 10^{-10}

```
> abs( 0.1*3 - 0.3 ) < 10^-10
[1] TRUE
```

To deal with this problem, there is a function called `all.equal()` that lets you test for equality but allows a small tolerance for rounding errors:

```
> all.equal( 0.1*3, 0.3 )
[1] TRUE
```

7.12.2 The recycling rule

There's one thing that I haven't mentioned about how vector arithmetic works in R, and that's the **recycling rule**. The easiest way to explain it is to give a simple example. Suppose I have two vectors of different length, `x` and `y`, and I want to add them together. It's not obvious what that actually means, so let's have a look at what R does:

```
> x <- c( 1,1,1,1,1,1 ) # x is length 6
> y <- c( 0,1 )          # y is length 2
> x + y                  # now add them:
[1] 1 2 1 2 1 2
```

As you can see from looking at this output, what R has done is "recycle" the value of the shorter vector (in this case `y`) several times. That is, the first element of `x` is added to the first element of `y`, and the second element of `x` is added to the second element of `y`. However, when R reaches the third element of `x` there isn't any corresponding element in `y`, so it returns to the beginning: thus, the third element of `x` is added to the *first* element of `y`. This process continues until R reaches the last element of `x`. And that's all there is to it really. The same recycling rule also applies for subtraction, multiplication and division. The only other thing I should note is that, if the length of the longer vector isn't an exact multiple of the length of the shorter one, R still does it, but also gives you a warning message:

```
> x <- c( 1,1,1,1,1 )      # x is length 5
> y <- c( 0,1 )            # y is length 2
> x + y                   # now add them:
```

²⁸Or at least, that's the default. If all your numbers are integers (whole numbers), then you can explicitly tell R to store them as integers by adding an `L` suffix at the end of the number. That is, an assignment like `x <- 2L` tells R to assign `x` a value of 2, and to store it as an integer rather than as a binary expansion. Type `?integer` for more details.

```
[1] 1 2 1 2 1
Warning message:
In x + y : longer object length is not a multiple of shorter object length
```

7.12.3 An introduction to environments

In this section I want to ask a slightly different question: what *is* the workspace exactly? This question seems simple, but there's a fair bit to it. This section can be skipped if you're not really interested in the technical details. In the description I gave earlier, I talked about the workspace as an abstract location in which R variables are stored. That's basically true, but it hides a couple of key details. For example, any time you have R open, it has to store *lots* of things in the computer's memory, not just your variables. For example, the `who()` function that I wrote has to be stored in memory somewhere, right? If it weren't I wouldn't be able to use it. That's pretty obvious. But equally obviously it's not in the workspace either, otherwise you should have seen it! Here's what's happening. R needs to keep track of a lot of different things, so what it does is organise them into **environments**, each of which can contain lots of different variables and functions. Your workspace is one such environment. Every package that you have loaded is another environment. And every time you call a function, R briefly creates a temporary environment in which the function itself can work, which is then deleted after the calculations are complete. So, when I type in `search()` at the command line

```
> search()
[1] ".GlobalEnv"      "package:lsr"      "tools:rstudio"
[4] "package:stats"   "package:graphics" "package:grDevices"
[7] "package:utils"    "package:datasets" "package:methods"
[10] "Autoloads"       "package:base"
```

what I'm actually looking at is a *sequence of environments*. The first one, `".GlobalEnv"` is the technically-correct name for your workspace (no-one really calls it that). And so when you type in `objects()` or `who()` what you're really doing is listing the contents of `".GlobalEnv"`. But there's no reason why we can't look up the contents of these other environments using the `objects()` function (currently `who()` doesn't support this). You just have to be a bit more explicit in your command. If I wanted to find out what is in the `package:stats` environment (i.e., the environment into which the contents of the `stats` package have been loaded), here's what I'd get

```
> objects("package:stats")
[1] "acf"                  "acf2AR"        "add.scope"
[4] "add1"                 "addmargins"    "aggregate"
[7] "aggregate.data.frame" "aggregate.default" "aggregate.ts"
[10] "AIC"                  "alias"         "anova"
[13] "anova.glm"           "anova.glmList" "anova.lm"
BLAH BLAH BLAH
```

where this time I've hidden a lot of output in the BLAH BLAH BLAH because the stats package contains about 500 functions. The key thing to understand then, is that you can access any of the R variables and functions that are stored in one of these environments, precisely because those are the environments that you have loaded!²⁹

²⁹For advanced users: that's a little over simplistic in two respects. First, it's a terribly imprecise way of talking about scoping. Second, it might give you the impression that all the variables in question are actually loaded into memory. That's not quite true, since that would be very wasteful of memory. Instead R has a "lazy loading" mechanism, in which what R actually does is create a "promise" to load those objects if they're actually needed. For details, check out the `delayedAssign()` function.

7.12.4 Attaching a data frame

The last thing I want to mention in this section is the `attach()` function, which you often see referred to in introductory R books. Whenever it is introduced, the author of the book usually mentions that the `attach()` function can be used to “attach” the data frame to the search path, so you don’t have to use the `$` operator. That is, if I use the command `attach(df)` to attach my data frame, I no longer need to type `df$variable`, and instead I can just type `variable`. This is true as far as it goes, but it’s very misleading and novice users often get led astray by this description, because it hides a lot of critical details.

Here is the very abridged description: when you use the `attach()` function, what R does is create an entirely new *environment* in the search path, just like when you load a package. Then, what it does is copy all of the variables in your data frame into this new environment. When you do this, however, you end up with two completely different versions of all your variables: one in the original data frame, and one in the new environment. Whenever you make a statement like `df$variable` you’re working with the variable inside the data frame; but when you just type `variable` you’re working with the copy in the new environment. And here’s the part that really upsets new users: *changes to one version are not reflected in the other version*. As a consequence, it’s really easy for R to end up with different value stored in the two different locations, and you end up really confused as a result.

To be fair to the writers of the `attach()` function, the help documentation does actually state all this quite explicitly, and they even give some examples of how this can cause confusion at the bottom of the help page. And I can actually see how it can be very useful to create copies of your data in a separate location (e.g., it lets you make all kinds of modifications and deletions to the data without having to touch the original data frame). However, I don’t think it’s helpful for new users, since it means you have to be very careful to keep track of which copy you’re talking about. As a consequence of all this, for the purpose of this book I’ve decided not to use the `attach()` function. It’s something that you can investigate yourself once you’re feeling a little more confidence with R, but I won’t do it here.

7.13 _____

Summary

Useful references for extending this are: [Spector \(2008\)](#) and [Teator \(2011\)](#).

8. Basic programming

Machine dreams hold a special vertigo.

– William Gibson

Up to this point in the book I've tried hard to avoid using the word "programming" because – at least in my experience – it's a word that can cause a lot of fear. For one reason or another, programming (like mathematics and statistics) is often perceived by people on the "outside" as a black art, a magical skill that can be learned only by some kind of super-nerd. I think this is a shame. It's certainly true that advanced programming is a very specialised skill: several different skills actually, since there's quite a lot of different kinds of programming out there. However, the *basics* of programming aren't all that hard, and you can accomplish a lot of very impressive things just using those basics.

With that in mind, the goal of this chapter is to discuss a few basic programming concepts and how to apply them in R. However, before I do, I want to make one further attempt to point out just how non-magical programming really is, via one very simple observation: *you already know how to do it*. Stripped to its essentials, programming is nothing more (and nothing less) than the process of writing out a bunch of instructions that a computer can understand. To phrase this slightly differently, when you write a computer program, you need to write it in a **programming language** that the computer knows how to interpret. R is one such language. Although I've been having you type all your commands at the command prompt, and all the commands in this book so far have been shown as if that's what I were doing, it's also quite possible (and as you'll see shortly, shockingly easy) to write a program using these R commands. In other words, if this is the first time reading this book, then you're only one short chapter away from being able to legitimately claim that you can program in R, albeit at a beginner's level.

8.1 _____

Scripts

Computer programs come in quite a few different forms: the kind of program that we're mostly interested in from the perspective of everyday data analysis using R is known as a **script**. The idea behind a script is that, instead of typing your commands into the R console one at a time, instead you write them all in a text file. Then, once you've finished writing them and saved the text file, you can get R to execute all the commands in your file by using the **source()** function. In a moment I'll show you exactly how this is done, but first I'd better explain why you should care.

8.1.1 Why use scripts?

Before discussing scripting and programming concepts in any more detail, it's worth stopping to ask why you should bother. After all, if you look at the R commands that I've used everywhere else this book, you'll notice that they're all formatted as if I were typing them at the command line. Outside this chapter you won't actually see any scripts. Do not be fooled by this. The reason that I've done it that way is purely for pedagogical reasons. My goal in this book is to teach statistics and to teach R. To that end, what *I've* needed to do is chop everything up into tiny little slices: each section tends to focus on one kind of statistical concept, and only a smallish number of R functions. As much as possible, I want you to see what each function does in isolation, one command at a time. By forcing myself to write everything as if it were being typed at the command line, it imposes a kind of discipline on me: it *prevents* me from piecing together lots of commands into one big script. From a teaching (and learning) perspective I think that's the right thing to do... but from a *data analysis* perspective, it is not. When you start analysing real world data sets, you will rapidly find yourself needing to write scripts.

To understand why scripts are so very useful, it may be helpful to consider the drawbacks to typing commands directly at the command prompt. The approach that we've been adopting so far, in which you type commands one at a time, and R sits there patiently in between commands, is referred to as the **interactive** style. Doing your data analysis this way is rather like having a conversation ... a very annoying conversation between you and your data set, in which you and the data aren't directly speaking to each other, and so you have to rely on R to pass messages back and forth. This approach makes a lot of sense when you're just trying out a few ideas: maybe you're trying to figure out what analyses are sensible for your data, or maybe just you're trying to remember how the various R functions work, so you're just typing in a few commands until you get the one you want. In other words, the interactive style is very useful as a tool for exploring your data. However, it has a number of drawbacks:

- *It's hard to save your work effectively.* You can save the workspace, so that later on you can load any variables you created. You can save your plots as images. And you can even save the history or copy the contents of the R console to a file. Taken together, all these things let you create a reasonably decent record of what you did. But it does leave a lot to be desired. It seems like you ought to be able to save a single file that R could use (in conjunction with your raw data files) and reproduce everything (or at least, everything interesting) that you did during your data analysis.
- *It's annoying to have to go back to the beginning when you make a mistake.* Suppose you've just spent the last two hours typing in commands. Over the course of this time you've created lots of new variables and run lots of analyses. Then suddenly you realise that there was a nasty typo in the first command you typed, so all of your later numbers are wrong. Now you have to fix that first command, and then spend another hour or so combing through the R history to try and recreate what you did.
- *You can't leave notes for yourself.* Sure, you can scribble down some notes on a piece of paper, or even save a Word document that summarises what you did. But what you really want to be able to do is write down an English translation of your R commands, preferably right "next to" the commands themselves. That way, you can look back at what you've done and actually remember what you were doing. In the simple exercises we've engaged in so far, it hasn't been all that hard to remember what you were doing or why you were doing it, but only because everything we've done could be done using only a few commands, and you've never been asked to reproduce your analysis six months after you originally did it! When your data analysis starts involving hundreds of variables, and requires quite complicated commands to work, then you really, really need to leave yourself some notes to explain your analysis to, well, yourself.
- *It's nearly impossible to reuse your analyses later, or adapt them to similar problems.* Suppose that, sometime in January, you are handed a difficult data analysis problem. After working on it

for ages, you figure out some really clever tricks that can be used to solve it. Then, in September, you get handed a really similar problem. You can sort of remember what you did, but not very well. You'd like to have a clean record of what you did last time, how you did it, and why you did it the way you did. Something like that would really help you solve this new problem.

- *It's hard to do anything except the basics.* There's a nasty side effect of these problems. Typos are inevitable. Even the best data analyst in the world makes a lot of mistakes. So the chance that you'll be able to string together dozens of correct R commands in a row are very small. So unless you have some way around this problem, you'll never really be able to do anything other than simple analyses.
- *It's difficult to share your work other people.* Because you don't have this nice clean record of what R commands were involved in your analysis, it's not easy to share your work with other people. Sure, you can send them all the data files you've saved, and your history and console logs, and even the little notes you wrote to yourself, but odds are pretty good that no-one else will really understand what's going on (trust me on this: I've been handed lots of random bits of output from people who've been analysing their data, and it makes very little sense unless you've got the original person who did the work sitting right next to you explaining what you're looking at)

Ideally, what you'd like to be able to do is something like this... Suppose you start out with a data set `myrawdata.csv`. What you want is a single document – let's call it `mydataanalysis.R` – that stores all of the commands that you've used in order to do your data analysis. Kind of similar to the R history but much more focused. It would only include the commands that you want to keep for later. Then, later on, instead of typing in all those commands again, you'd just tell R to run all of the commands that are stored in `mydataanalysis.R`. Also, in order to help you make sense of all those commands, what you'd want is the ability to add some notes or *comments* within the file, so that anyone reading the document for themselves would be able to understand what each of the commands actually does. But these comments wouldn't get in the way: when you try to get R to run `mydataanalysis.R` it would be smart enough to recognise that these comments are for the benefit of humans, and so it would ignore them. Later on you could tweak a few of the commands inside the file (maybe in a new file called `mynewdatanalysis.R`) so that you can adapt an old analysis to be able to handle a new problem. And you could email your friends and colleagues a copy of this file so that they can reproduce your analysis themselves.

In other words, what you want is a *script*.

8.1.2 Our first script

Okay then. Since scripts are so terribly awesome, let's write one. To do this, open up a simple text editing program, like TextEdit (on a Mac) or Notebook (on Windows). Don't use a fancy word processing program like Microsoft Word or OpenOffice: use the simplest program you can find. Open a new text document, and type some R commands, hitting enter after each command. Let's try using `x <- "hello world"` and `print(x)` as our commands. Then save the document as `hello.R`, and remember to save it as a plain text file: don't save it as a word document or a rich text file. Just a boring old plain text file. Also, when it asks you *where* to save the file, save it to whatever folder you're using as your working directory in R. At this point, you should be looking at something like Figure 8.1. And if so, you have now successfully written your first R program. Because I don't want to take screenshots for every single script, I'm going to present scripts using extracts formatted as follows:

```
hello.R
1   x <- "hello world"
2   print(x)
```

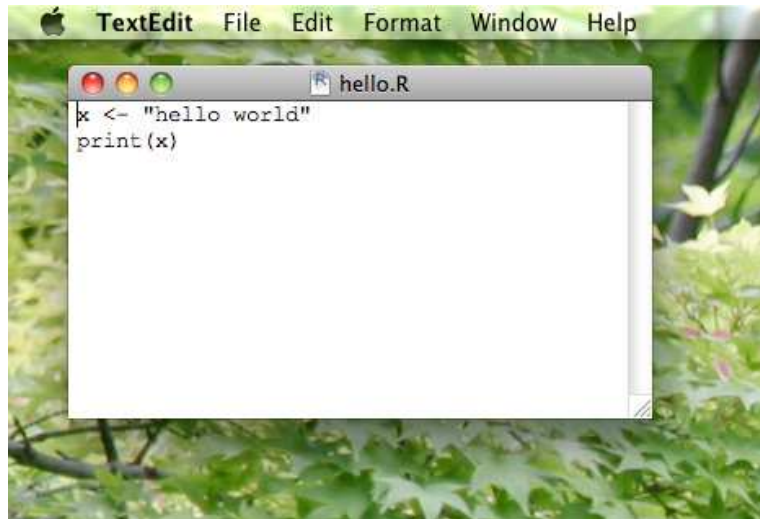


Figure 8.1: A screenshot showing the `hello.R` script if you open it in the default text editor (TextEdit) on a Mac. Using a simple text editor like TextEdit on a Mac or Notepad on Windows isn't actually the best way to write your scripts, but it is the simplest. More to the point, it highlights the fact that a script really is just an ordinary text file.

.....

The line at the top is the filename, and not part of the script itself. Below that, you can see the two R commands that make up the script itself. Next to each command I've included the line numbers. You don't actually type these into your script, but a lot of text editors (including the one built into Rstudio that I'll show you in a moment) will show line numbers, since it's a very useful convention that allows you to say things like “line 1 of the script creates a new variable, and line 2 prints it out”.

So how do we run the script? Assuming that the `hello.R` file has been saved to your working directory, then you can run the script using the following command:

```
> source( "hello.R" )
```

If the script file is saved in a different directory, then you need to specify the path to the file, in exactly the same way that you would have to when loading a data file using `load()`. In any case, when you type this command, R opens up the script file: it then reads each command in the file in the same order that they appear in the file, and executes those commands in that order. The simple script that I've shown above contains two commands. The first one creates a variable `x` and the second one prints it on screen. So, when we run the script, this is what we see on screen:

```
> source( "hello.R" )
[1] "hello world"
```

If we inspect the workspace using a command like `who()` or `objects()`, we discover that R has created the new variable `x` within the workspace, and not surprisingly `x` is a character string containing the text `"hello world"`. And just like that, you've written your first program R. It really is that simple.

8.1.3 Using Rstudio to write scripts

In the example above I assumed that you were writing your scripts using a simple text editor. However, it's usually more convenient to use a text editor that is specifically designed to help you write scripts. There's a lot

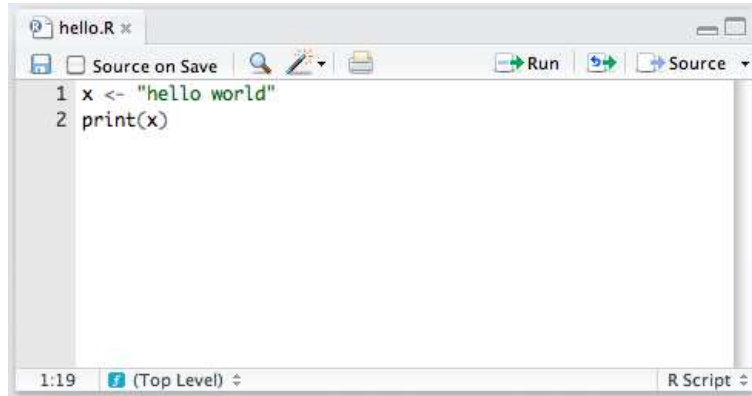


Figure 8.2: A screenshot showing the `hello.R` script open in Rstudio. Assuming that you’re looking at this document in colour, you’ll notice that the “hello world” text is shown in green. This isn’t something that you do yourself: that’s Rstudio being helpful. Because the text editor in Rstudio “knows” something about how R commands work, it will highlight different parts of your script in different colours. This is useful, but it’s not actually part of the script itself.

of these out there, and experienced programmers will all have their own personal favourites. For our purposes, however, we can just use the one built into Rstudio. To create new script file in R studio, go to the “File ” menu, select the “New” option, and then click on “R script”. This will open a new window within the “source” panel. Then you can type the commands you want (or **code** as it is generally called when you’re typing the commands into a script file) and save it when you’re done. The nice thing about using Rstudio to do this is that it automatically changes the colour of the text to indicate which parts of the code are comments and which are parts are actual R commands (these colours are called **syntax highlighting**, but they’re not actually part of the file – it’s just Rstudio trying to be helpful. To see an example of this, let’s open up our `hello.R` script in Rstudio. To do this, go to the “File” menu again, and select “Open...”. Once you’ve opened the file, you should be looking at something like Figure 8.2. As you can see (if you’re looking at this book in colour) the character string “hello world” is highlighted in green.

Using Rstudio for your text editor is convenient for other reasons too. Notice in the top right hand corner of Figure 8.2 there’s a little button that reads “Source”? If you click on that, Rstudio will construct the relevant `source()` command for you, and send it straight to the R console. So you don’t even have to type in the `source()` command, which actually I think is a great thing, because it really bugs me having to type all those extra keystrokes every time I want to run my script. Anyway, Rstudio provide several other convenient little tools to help make scripting easier, but I won’t discuss them here.

8.1.4 Commenting your script

When writing up your data analysis as a script, one thing that is generally a good idea is to include a lot of comments in the code. That way, if someone else tries to read it (or if you come back to it several days, weeks, months or years later) they can figure out what’s going on. As a beginner, I think it’s especially useful to comment thoroughly, partly because it gets you into the habit of commenting the code, and partly because the simple act of typing in an explanation of what the code does will help you keep it clear in your own mind what you’re trying to achieve. To illustrate this idea, consider the following script:

```
itngscript.R
1 # A script to analyse nightgarden.Rdata
2 # author: Dan Navarro
3 # date: 22/11/2011
```

```

4
5   # Load the data, and tell the user that this is what we're
6   # doing. Note that this assumes the nightgarden data file
7   # is in the working directory.
8   cat( "loading data from nightgarden.Rdata...\n" )
9   load( "nightgarden.Rdata" )

10
11  # Create a cross tabulation and print it out:
12  cat( "tabulating data...\n" )
13  itng.table <- table( speaker, utterance )
14  print( itng.table )

```

Firstly, as you can see from this extract I've done a tiny bit of syntax highlighting here, greying out the comments a bit so that you can visually distinguish between the commands that R will execute and the comments which it will ignore. Secondly, notice that I've gone a bit overboard with my commenting: at the top of the script I've explained the purpose of the script, who wrote it, and when it was written. Then, throughout the script file itself I've added a lot of comments explaining what each section of the code actually does. In real life people don't tend to comment this thoroughly, but the basic idea is a very good one: you really do want your script to explain itself. Nevertheless, as you'd expect R completely ignores all of the commented parts. When we run this script, this is what we see on screen:

```

> source( "itngscript.R" )
loading data from nightgarden.Rdata...
tabulating data...
      utterance
speaker    ee onk oo pip
  makka-pakka  0  2  0  2
  tombliboo    1  0  1  0
  upsy-daisy   0  2  0  2

```

Even here, notice that the script announces its behaviour. The first two lines of the output tell us a lot about what the script is actually doing behind the scenes (the code do to this corresponds to the two `cat()` commands on lines 8 and 12 of the script). It's usually a pretty good idea to do this, since it helps ensure that the output makes sense when the script is executed.

8.1.5 Differences between scripts and the command line

For the most part, commands that you insert into a script behave in exactly the same way as they would if you typed the same thing in at the command line. The one major exception to this is that if you want a variable to be printed on screen, you need to explicitly tell R to print it. You can't just type the name of the variable. For example, our original `hello.R` script produced visible output. The following script does not:

```

silenthello.R
1   x <- "hello world"
2   x

```

It *does* still create the variable `x` when you `source()` the script, but it won't print anything on screen.

However, apart from the fact that scripts don't use "auto-printing" as it's called, there aren't a lot of differences in the underlying mechanics. There are a few stylistic differences though. For instance, if you want to load a package at the command line, you would generally use the `library()` function. If you want do to it from a script, it's conventional to use `require()` instead. The two commands are basically identical, the only difference being that if the package doesn't exist, `require()` produces a warning whereas `library()` gives you an error. Stylistically, what this means is that if the `require()` command fails in your script, R will boldly continue on and try to execute the rest of the script. Often that's what you'd like to see happen, so it's better to use `require()`. Clearly, however, you can get by just fine using the `library()` command for everyday usage.

8.1.6 Done!

At this point, you've learned the basics of scripting. You are now officially allowed to say that you can program in R, though you probably shouldn't say it too loudly. There's a *lot* more to learn, but nevertheless, if you can write scripts like these then what you are doing is in fact basic programming. The rest of this chapter is devoted to introducing some of the key commands that you need in order to make your programs more powerful; and to help you get used to thinking in terms of scripts, for the rest of this chapter I'll write up most of my extracts as scripts.

8.2

Loops

The description I gave earlier for how a script works was a tiny bit of a lie. Specifically, it's not necessarily the case that R starts at the top of the file and runs straight through to the end of the file. For all the scripts that we've seen so far that's exactly what happens, and unless you insert some commands to explicitly alter how the script runs, that is what will *always* happen. However, you actually have quite a lot of flexibility in this respect. Depending on how you write the script, you can have R repeat several commands, or skip over different commands, and so on. This topic is referred to as **flow control**, and the first concept to discuss in this respect is the idea of a **loop**. The basic idea is very simple: a loop is a block of code (i.e., a sequence of commands) that R will execute over and over again until some termination criterion is met. Looping is a very powerful idea. There are three different ways to construct a loop in R, based on the **while**, **for** and **repeat** functions. I'll only discuss the first two in this book.

8.2.1 The **while** loop

A **while** loop is a simple thing. The basic format of the loop looks like this:

```
while ( CONDITION ) {  
    STATEMENT1  
    STATEMENT2  
    ETC  
}
```

The code corresponding to CONDITION needs to produce a logical value, either **TRUE** or **FALSE**. Whenever R encounters a **while** statement, it checks to see if the CONDITION is **TRUE**. If it is, then R goes on to execute all of the commands inside the curly brackets, proceeding from top to bottom as usual. However, when it gets to the bottom of those statements, it moves back up to the **while** statement. Then, like the mindless automaton it is, it checks to see if the CONDITION is **TRUE**. If it is, then R goes on to execute all ... well, you get the idea. This continues endlessly until at some point the CONDITION turns out to be **FALSE**. Once that happens, R jumps to the bottom of the loop (i.e., to the **}** character), and then continues on with whatever commands appear next in the script.

To start with, let's keep things simple, and use a **while** loop to calculate the smallest multiple of 17 that is greater than or equal to 1000. This is a very silly example since you can actually calculate it using simple arithmetic operations, but the point here isn't to do something novel. The point is to show how to write a **while** loop. Here's the script:

whileexample.R

```
1   x <- 0  
2   while ( x < 1000 ) {  
3       x <- x + 17
```

```

4      }
5  print( x )

```

When we run this script, R starts at the top and creates a new variable called `x` and assigns it a value of 0. It then moves down to the loop, and “notices” that the condition here is `x < 1000`. Since the current value of `x` is zero, the condition is true, so it enters the body of the loop (inside the curly braces). There’s only one command here¹ which instructs R to increase the value of `x` by 17. R then returns to the top of the loop, and rechecks the condition. The value of `x` is now 17, but that’s still less than 1000, so the loop continues. This cycle will continue for a total of 59 iterations, until finally `x` reaches a value of 1003 (i.e., $59 \times 17 = 1003$). At this point, the loop stops, and R finally reaches line 5 of the script, prints out the value of `x` on screen, and then halts. Let’s watch:

```

> source( "whileexample.R" )
[1] 1003

```

Truly fascinating stuff.

8.2.2 The `for` loop

The `for` loop is also pretty simple, though not quite as simple as the `while` loop. The basic format of this loop goes like this:

```

for ( VAR in VECTOR ) {
  STATEMENT1
  STATEMENT2
  ETC
}

```

In a `for` loop, R runs a fixed number of iterations. We have a VECTOR which has several elements, each one corresponding to a possible value of the variable VAR. In the first iteration of the loop, VAR is given a value corresponding to the first element of VECTOR; in the second iteration of the loop VAR gets a value corresponding to the second value in VECTOR; and so on. Once we’ve exhausted all of the values in VECTOR, the loop terminates and the flow of the program continues down the script.

Once again, let’s use some very simple examples. Firstly, here is a program that just prints out the word “hello” three times and then stops:

```

forexample.R

1  for ( i in 1:3 ) {
2    print( "hello" )
3  }

```

This is the simplest example of a `for` loop. The vector of possible values for the `i` variable just corresponds to the numbers from 1 to 3. Not only that, the body of the loop doesn’t actually depend on `i` at all. Not surprisingly, here’s what happens when we run it:

```

> source( "forexample.R" )
[1] "hello"
[1] "hello"
[1] "hello"

```

However, there’s nothing that stops you from using something non-numeric as the vector of possible values, as the following example illustrates. This time around, we’ll use a character vector to control our loop, which in this

¹As an aside: if there’s only a single command that you want to include inside your loop, then you don’t actually need to bother including the curly braces at all. However, until you’re comfortable programming in R I’d advise *always* using them, even when you don’t have to.

case will be a vector of `words`. And what we'll do in the loop is get R to convert the word to upper case letters, calculate the length of the word, and print it out. Here's the script:

```
forexample2.R
1   #the words
2   words <- c("it", "was", "the", "dirty", "end", "of", "winter")
3
4   #loop over the words
5   for ( w in words ) {
6
7     w.length <- nchar( w )      # calculate the number of letters
8     W <- toupper( w )          # convert the word to upper case letters
9     msg <- paste( W, "has", w.length, "letters" )    # a message to print
10    print( msg )                # print it
11
12 }
```

And here's the output:

```
> source( "forexample2.R" )
[1] "IT has 2 letters"
[1] "WAS has 3 letters"
[1] "THE has 3 letters"
[1] "DIRTY has 5 letters"
[1] "END has 3 letters"
[1] "OF has 2 letters"
[1] "WINTER has 6 letters"
```

Again, pretty straightforward I hope.

8.2.3 A more realistic example of a loop

To give you a sense of how you can use a loop in a more complex situation, let's write a simple script to simulate the progression of a mortgage. Suppose we have a nice young couple who borrow \$300000 from the bank, at an annual interest rate of 5%. The mortgage is a 30 year loan, so they need to pay it off within 360 months total. Our happy couple decide to set their monthly mortgage payment at \$1600 per month. Will they pay off the loan in time or not? Only time will tell.² Or, alternatively, we could simulate the whole process and get R to tell us. The script to run this is a fair bit more complicated.

```
mortgage.R
1   # set up
2   month <- 0  # count the number of months
3   balance <- 300000 # initial mortgage balance
4   payments <- 1600 # monthly payments
5   interest <- 0.05 # 5% interest rate per year
6   total.paid <- 0 # track what you've paid the bank
7
8   # convert annual interest to a monthly multiplier
9   monthly.multiplier <- (1+interest) ^ (1/12)
10
```

²Okay, fine. This example is still a bit ridiculous, in three respects. Firstly, the bank absolutely will not let the couple pay less than the amount required to terminate the loan in 30 years. Secondly, a constant interest rate of 30 years is hilarious. Thirdly, you can solve this much more efficiently than through brute force simulation. However, we're not exactly in the business of being realistic or efficient here.

```

11     # keep looping until the loan is paid off...
12     while ( balance > 0 ) {
13
14         # do the calculations for this month
15         month <- month + 1 # one more month
16         balance <- balance * monthly.multiplier # add the interest
17         balance <- balance - payments # make the payments
18         total.paid <- total.paid + payments # track the total paid
19
20         # print the results on screen
21         cat( "month", month, ": balance", round(balance), "\n" )
22
23     } # end of loop
24
25     # print the total payments at the end
26     cat("total payments made", total.paid, "\n" )

```

To explain what's going on, let's go through it carefully. In the first block of code (lines 2-6) all we're doing is specifying all the variables that define the problem. The loan starts with a `balance` of \$300,000 owed to the bank on `month` zero, and at that point in time the `total.paid` money is nothing. The couple is making monthly `payments` of \$1600, at an annual `interest` rate of 5%. Next, on line 9, we convert the annual percentage interest into a monthly multiplier. That is, the number that you have to multiply the current balance by each month in order to produce an annual interest rate of 5%. An annual interest rate of 5% implies that, if no payments were made over 12 months the balance would end up being 1.05 times what it was originally, so the *annual* multiplier is 1.05. To calculate the monthly multiplier, we need to calculate the 12th root of 1.05 (i.e., raise 1.05 to the power of 1/12). We store this value in as the `monthly.multiplier` variable, which as it happens corresponds to a value of about 1.004. All of which is a rather long winded way of saying that the *annual* interest rate of 5% corresponds to a *monthly* interest rate of about 0.4%.

Anyway... all of that is really just setting the stage. It's not the interesting part of the script. The interesting part (such as it is) is the loop which unfolds over lines 12-23. The `while` statement on line 12 tells R that it needs to keep looping until the `balance` reaches zero (or less, since it might be that the final payment of \$1600 pushes the balance below zero). Then, inside the body of the loop, we have two different blocks of code. In the first bit, on lines 15-18, we do all the number crunching. Firstly we increase the value `month` by 1 (on line 15). Next, the bank charges the interest (on line 16), so the `balance` goes up. Then, the couple makes their monthly payment (on line 17) and the `balance` goes down. Finally, we keep track of the total amount of money that the couple has paid so far, by adding the `payments` to the running tally (on line 18). After having done all this number crunching, we tell R (on line 21) to issue the couple with a very terse monthly statement, which just indicates how many months they've been paying the loan and how much money they still owe the bank. Which is rather rude of us really. I've grown attached to this couple and I really feel they deserve better than that. But, that's banks for you.

In any case, the key thing here is the tension between the increase in `balance` on line 16 and the decrease on line 17. As long as the decrease is bigger, then the balance will eventually drop to zero and the loop will eventually terminate. If not, the loop will continue forever! This is actually very bad programming on my part: I really should have included something to force R to stop if this goes on too long. However, I haven't shown you how to evaluate "if" statements yet, so we'll just have to hope that the author of the book has rigged the example so that the code actually runs. Hm. I wonder what the odds of that are? Anyway, assuming that the loop does eventually terminate, there's one last line of code (line 26) that prints out the total amount of money that the couple handed over to the bank over the lifetime of the loan.

Now that I've explained everything in the script in tedious detail, let's run it and see what happens:

```

> source( "mortgage.R" )
month 1 : balance 299622
month 2 : balance 299243
month 3 : balance 298862
month 4 : balance 298480
month 5 : balance 298096

```

```
BLAH BLAH BLAH
```

```
month 352 : balance 4806
month 353 : balance 3226
month 354 : balance 1639
month 355 : balance 46
month 356 : balance -1554
total payments made 569600
```

So our nice young couple have paid off their \$300,000 loan in just 4 months shy of the 30 year term of their loan, at a bargain basement price of \$568,046 (since $569600 - 1554 = 568046$). A happy ending!

8.3 _____

Conditional statements

A second kind of flow control that programming languages provide is the ability to evaluate **conditional statements**. Unlike loops, which can repeat over and over again, a conditional statement only executes once, but it can switch between different possible commands depending on a CONDITION that is specified by the programmer. The power of these commands is that they allow the program itself to make choices, and in particular, to make different choices depending on the context in which the program is run. The most prominent example of a conditional statement is the **if** statement, and the accompanying **else** statement. The basic format of an **if** statement in R is as follows:

```
if ( CONDITION ) {
  STATEMENT1
  STATEMENT2
  ETC
}
```

And the execution of the statement is pretty straightforward. If the CONDITION is true, then R will execute the statements contained in the curly braces. If the CONDITION is false, then it does not. If you want to, you can extend the **if** statement to include an **else** statement as well, leading to the following syntax:

```
if ( CONDITION ) {
  STATEMENT1
  STATEMENT2
  ETC
} else {
  STATEMENT3
  STATEMENT4
  ETC
}
```

As you'd expect, the interpretation of this version is similar. If the CONDITION is true, then the contents of the first block of code (i.e., STATEMENT1, STATEMENT2, ETC) are executed; but if it is false, then the contents of the second block of code (i.e., STATEMENT3, STATEMENT4, ETC) are executed instead.

To give you a feel for how you can use **if** and **else** to do something useful, the example that I'll show you is a script that prints out a different message depending on what day of the week you run it. We can do this making use of some of the tools that we discussed in Section 7.11.3. Here's the script:

[ifelseexample.R](#)

```

1   # find out what day it is...
2   today <- Sys.Date()      # pull the date from the system clock
3   day <- weekdays( today ) # what day of the week it is
4
5   # now make a choice depending on the day...
6   if ( day == "Monday" ) {
7     print( "I don't like Mondays" )
8   } else {
9     print( "I'm a happy little automaton" )
10 }

```

Since today happens to be a Tuesday, when I run the script here's what happens:

```

> source( "ifelseexample.R" )
[1] "I'm a happy little automaton"

```

There are other ways of making conditional statements in R. In particular, the `ifelse()` function and the `switch()` functions can be very useful in different contexts. However, my main aim in this chapter is to briefly cover the very basics, so I'll move on.

8.4 _____

Writing functions

In this section I want to talk about functions again. Functions were introduced in Section 3.5, but you've learned a lot about R since then, so we can talk about them in more detail. In particular, I want to show you how to create your own. To stick with the same basic framework that I used to describe loops and conditionals, here's the syntax that you use to create a function:

```

FNAME <- function ( ARG1, ARG2, ETC ) {
  STATEMENT1
  STATEMENT2
  ETC
  return( VALUE )
}

```

What this does is create a function with the name `FNAME`, which has arguments `ARG1`, `ARG2` and so forth. Whenever the function is called, R executes the statements in the curly braces, and then outputs the contents of `VALUE` to the user. Note, however, that R does not execute the commands inside the function in the workspace. Instead, what it does is create a temporary local environment: all the internal statements in the body of the function are executed there, so they remain invisible to the user. Only the final results in the `VALUE` are returned to the workspace.

To give a simple example of this, let's create a function called `quadruple()` which multiplies its inputs by four. In keeping with the approach taken in the rest of the chapter, I'll use a script to do this:

```

functionexample.R

1   quadruple <- function(x) {
2     y <- x*4
3     return(y)
4   }

```

When we run this script, as follows

```

> source( "functionexample.R" )

```

nothing appears to have happened, but there is a new object created in the workspace called `quadruple`. Not surprisingly, if we ask R to tell us what kind of object it is, it tells us that it is a function:

```
> class( quadruple )
[1] "function"
```

And now that we've created the `quadruple()` function, we can call it just like any other function. And if I want to store the output as a variable, I can do this:

```
> my.var <- quadruple(10)
> print(my.var)
[1] 40
```

An important thing to recognise here is that the two internal variables that the `quadruple()` function makes use of, `x` and `y`, stay internal. That is, if we inspect the contents of the workspace,

```
> who()
--- Name --- --- Class ---
my.var      numeric
quadruple   function
```

we see only the `quadruple()` function itself, as well as the `my.var` variable that we just created.

Now that we know how to create our own functions in R, it's probably a good idea to talk a little more about some of the other properties of functions that I've been glossing over. To start with, let's take this opportunity to type the name of the function at the command line without the parentheses:

```
> quadruple
function(x)  y <- x*4  return(y)
```

As you can see, when you type the name of a function at the command line, R prints out the underlying source code that we used to define the function in the first place. In the case of the `quadruple()` function, this is quite helpful to us – we can read this code and actually see what the function does. For other functions, this is less helpful, as we saw back in Section 3.5 when we tried typing `citation` rather than `citation()`.

8.4.1 Function arguments revisited

Okay, now that we are starting to get a sense for how functions are constructed, let's have a look at two, slightly more complicated functions that I've created. The source code for these functions is contained within the `functionexample2.R` and `functionexample3.R` scripts. Let's start by looking at the first one:

```
functionexample2.R
1  pow <- function( x, y = 1 ) {
2    out <- x^y # raise x to the power y
3    return( out )
4 }
```

and if we type `source("functionexample2.R")` to load the `pow()` function into our workspace, then we can make use of it. As you can see from looking at the code for this function, it has two arguments `x` and `y`, and all it does is raise `x` to the power of `y`. For instance, this command

```
> pow( x=3, y=2 )
[1] 9
```

calculates the value of 3^2 . The interesting thing about this function isn't what it does, since R already has perfectly good mechanisms for calculating powers. Rather, notice that when I defined the function, I specified `y=1` when listing the arguments? That's the default value for `y`. So if we enter a command without specifying a value for `y`, then the function assumes that we want `y=1`:

```
> pow( x=3 )
[1] 3
```

However, since I didn't specify any default value for `x` when I defined the `pow()` function, we always need to input a value for `x`. If we don't R will spit out an error message.

So now you know how to specify default values for an argument. The other thing I should point out while I'm on this topic is the use of the `...` argument. The `...` argument is a special construct in R which is only used within functions. It is used as a way of matching against multiple user inputs: in other words, `...` is used as a mechanism to allow the user to enter as many inputs as they like. I won't talk at all about the low-level details of how this works at all, but I will show you a simple example of a function that makes use of it. To that end, consider the following script:

`functionexample3.R`

```
1  doubleMax <- function( ... ) {
2      max.val <- max( ... )    # find the largest value in ...
3      out <- 2 * max.val     # double it
4      return( out )
5 }
```

When we type `source("functionexample3.R")`, R creates the `doubleMax()` function. You can type in as many inputs as you like. The `doubleMax()` function identifies the largest value in the inputs, by passing all the user inputs to the `max()` function, and then doubles it. For example:

```
> doubleMax( 1,2,5 )
[1] 10
```

8.4.2 There's more to functions than this

There's a lot of other details to functions that I've hidden in my description in this chapter. Experienced programmers will wonder exactly how the "scoping rules" work in R,³ or want to know how to use a function to create variables in other environments⁴, or if function objects can be assigned as elements of a list⁵ and probably hundreds of other things besides. However, I don't want to have this discussion get too cluttered with details, so I think it's best – at least for the purposes of the current book – to stop here.

8.5

Implicit loops

There's one last topic I want to discuss in this chapter. In addition to providing the explicit looping structures via `while` and `for`, R also provides a collection of functions for **implicit loops**. What I mean by this is that these are functions that carry out operations very similar to those that you'd normally use a loop for. However, instead of typing out the whole loop, the whole thing is done with a single command. The main reason why this can be handy is that – due to the way that R is written – these implicit looping functions are usually about to do the same calculations much faster than the corresponding explicit loops. In most applications that beginners might want to undertake, this probably isn't very important, since most beginners tend to start out working with fairly small data sets and don't usually need to undertake extremely time consuming number crunching. However, because you often see these functions referred to in other contexts, it may be useful to very briefly discuss a few of them.

³Lexical scope.

⁴The `assign()` function.

⁵Yes.

The first and simplest of these functions is `sapply()`. The two most important arguments to this function are `X`, which specifies a vector containing the data, and `FUN`, which specifies the name of a function that should be applied to each element of the data vector. The following example illustrates the basics of how it works:

```
> words <- c("along", "the", "loom", "of", "the", "land")
> sapply( X = words, FUN = nchar )
along   the   loom   of   the   land
      5     3     4     2     3     4
```

Notice how similar this is to the second example of a `for` loop in Section 8.2.2. The `sapply()` function has implicitly looped over the elements of `words`, and for each such element applied the `nchar()` function to calculate the number of letters in the corresponding word.

The second of these functions is `tapply()`, which has three key arguments. As before `X` specifies the data, and `FUN` specifies a function. However, there is also an `INDEX` argument which specifies a grouping variable.⁶ What the `tapply()` function does is loop over all of the different values that appear in the `INDEX` variable. Each such value defines a group: the `tapply()` function constructs the subset of `X` that corresponds to that group, and then applies the function `FUN` to that subset of the data. This probably sounds a little abstract, so let's consider a specific example, using the `nightgarden.Rdata` file that we used in Chapter 7.

```
> gender <- c( "male", "male", "female", "female", "male" )
> age <- c( 10, 12, 9, 11, 13 )
> tapply( X = age, INDEX = gender, FUN = mean )
    female      male
10.00000 11.66667
```

In this extract, what we're doing is using `gender` to define two different groups of people, and using their `ages` as the data. We then calculate the `mean()` of the ages, separately for the males and the females. A closely related function is `by()`. It actually does the same thing as `tapply()`, but the output is formatted a bit differently. This time around the three arguments are called `data`, `INDICES` and `FUN`, but they're pretty much the same thing. An example of how to use the `by()` function is shown in the following extract:

```
> by( data = age, INDICES = gender, FUN = mean )
gender: female
[1] 10
-----
gender: male
[1] 11.66667
```

The `tapply()` and `by()` functions are quite handy things to know about, and are pretty widely used. However, although I do make passing reference to the `tapply()` later on, I don't make much use of them in this book.

Before moving on, I should mention that there are several other functions that work along similar lines, and have suspiciously similar names: `lapply`, `mapply`, `apply`, `vapply`, `rapply` and `eapply`. However, none of these come up anywhere else in this book, so all I wanted to do here is draw your attention to the fact that they exist.

8.6

Summary

- Writing and using scripts (Section 8.1).
- Using loops (Section 8.2) and implicit loops (Section 8.5).
- Making conditional statements (Section 8.3)
- Writing your own functions (Section 8.4)

⁶Or a list of such variables.

Part IV.

Statistical theory

9. Introduction to probability

[God] has afforded us only the twilight . . . of Probability.

– John Locke

Up to this point, we've discussed some of the key ideas in experimental design, and we've talked a little about how you can summarise a data set. To a lot of people, this is all there is to statistics: it's about calculating averages, collecting all the numbers, drawing pictures, and putting them all in a report somewhere. Kind of like stamp collecting, but with numbers. However, statistics covers much more than that. In fact, descriptive statistics is one of the smallest parts of statistics, and one of the least powerful. The bigger and more useful part of statistics is that it provides that let you make inferences about data.

Once you start thinking about statistics in these terms – that statistics is there to help us draw inferences from data – you start seeing examples of it everywhere. For instance, here's a tiny extract from a newspaper article in the Sydney Morning Herald (30 Oct 2010):

"I have a tough job," the Premier said in response to a poll which found her government is now the most unpopular Labor administration in polling history, with a primary vote of just 23 per cent.

This kind of remark is entirely unremarkable in the papers or in everyday life, but let's have a think about what it entails. A polling company has conducted a survey, usually a pretty big one because they can afford it. I'm too lazy to track down the original survey, so let's just imagine that they called 1000 NSW voters at random, and 230 (23%) of those claimed that they intended to vote for the ALP. As of the 2010 Federal election, the Australian Electoral Commission reports that NSW has 4,610,795 enrolled voters; so the opinions of the remaining 4,609,795 voters (about 99.98% of voters) remain unknown to us. Even assuming that no-one lied to the polling company (an entirely unreasonable assumption, I should add) the only thing we can say with absolute 100% confidence is that the true ALP primary vote is somewhere between 230/4610795 (about 0.005%) and 4610025/4610795 (about 99.83%). So, on what basis is it legitimate for the polling company, the newspaper, and the readership to conclude that the ALP primary vote is only about 23%?

The answer to the question is pretty obvious: if I call 1000 people at random, and 230 of them say they intend to vote for the ALP, then it seems very unlikely that these are the *only* 230 people out of the entire voting public who actually intend to do so. In other words, we assume that the data collected by the polling company is pretty representative of the population at large. But how representative? Would we be surprised to discover that the true ALP primary vote is actually 24%? 29%? 37%? At this point everyday intuition starts to break down a bit. No-one would be surprised by 24%, and everybody would be surprised by 37%, but it's a bit hard to say whether 29% is plausible. We need some more powerful tools than just looking at the numbers and guessing.

Inferential statistics provides the tools that we need to answer these sorts of questions, and since these kinds of questions lie at the heart of the scientific enterprise, they take up the lions share of every introductory course on statistics and research methods. However, the theory of statistical inference is

built on top of **probability theory**. And it is to probability theory that we must now turn. This discussion of probability theory is basically background: there's not a lot of statistics per se in this chapter, and you don't need to understand this material in as much depth as the other chapters in this part of the book. Nevertheless, because probability theory does underpin so much of statistics, it's worth covering some of the basics. In short, you can "get away" with a much more cursory reading of this chapter than you can for most of the other chapters in the book, but you should still read it!

9.1

Probability theory v. statistical inference

Probability theory and inferential statistics are closely related to one another, but they're not quite the same. Statistics is fundamentally about data analysis, whereas probability theory is a bit more abstract: it's a branch of maths, really. We're not going to discuss the mathematics in a lot of detail in this book, but here's some simple examples of the sorts of things that probability theory is concerned with:

- What are the chances of a fair coin coming up heads 10 times in a row?
- How likely is it that five cards drawn from a perfectly shuffled deck will all be hearts?

In contrast, here's some questions that are more statistical in nature:

- If a coin comes up heads 10 times in a row, how likely is it that the coin is fair?
- If five cards off the top of the deck are all hearts, how likely is it that the deck was properly shuffled?

Notice the key difference between the two types of question. The probabilistic ones *assume* that we can write down a "probability model" of some kind, and then calculate "the chances" of an event. In contrast, the statistical questions tend to go the other way: they assume that we have observed some event, and now we have to figure out what "probability model" generated the data. For instance, consider the "coin flipping" example, the probabilistic question stated at the outset that the coin was fair. So we know that the probability that any given coin comes up heads, which we could write $P(H)$, is fixed at 50%. Or, to be a bit more precise, we'd write

$$P(H) = 0.5$$

(as we'll see later, in the same way that percentages go from 0% to 100%, probabilities go from 0 to 1). This knowledge, this statement that $P(H) = 0.5$, is a simple example of a **model**: that is, a precise, mathematically-phrased theory about the data and how it was generated. Since we know what the probability of a single head is, we can use probability theory to calculate the probability of 10 heads.

Now think about the statistical version of the problem. What we have actually observed is a data set, a "sample" if you will, so we'll again refer to that as X (i.e., X is the set of 10 heads that we observed). What we suspect is that these data are being generated from a probabilistic model very similar to the one discussed above, where there is some probability $P(H)$ that the coin comes up heads, but we actually don't know if the coin is fair or not. In other words, we have an unknown **parameter** that needs to be **estimated**. Let's denote this parameter by θ (Greek letter theta), and so we can write

$$\theta = P(H)$$

From a statistics perspective, our goal is to use the data X to figure out what the true value of θ is. Notice how this contrasts with the probabilistic question, in which we use θ to figure out how likely X is. The two questions run in opposite directions to each other, though the probabilistic question is much easier to solve than the statistical one. In the next chapter, I'll talk about how to solve this estimation problem, but I hope it's obvious that whatever the answer to the estimation problem is, it's obviously going to rely on probability theory. So that's where we need to begin.

Basic probability theory

It might seem like the obvious place to start would be to try to define “probability”. However, as I’ll discuss in the next section, this is actually quite difficult, and is a source of real disagreement among statisticians. So, instead, I’ll just start by giving a very trite definition... for now, let’s just agree that the probability of an event is the “chance” that the event will happen. Obviously this is completely tautological (defining probability in terms of chance), but since everyone already has an intuitive feeling for what “chance” means, it’ll do for now. With that in mind, instead of trying to say what probability *is*, I’ll spend this section talking about the *rules* that probabilities must obey, since everyone at least agrees on those. And in order to do so, I’m going to have to talk about my pants.

9.2.1 Introducing probability distributions

One of the disturbing truths about my life is that I only own 5 pairs of pants: three pairs of jeans, the bottom half of a suit, and a pair of tracksuit pants. Even sadder, I’ve given them names: I call them X_1 , X_2 , X_3 , X_4 and X_5 . I really do: that’s why they call me Mister Imaginative. Now, on any given day, I pick out exactly one of pair of pants to wear. Not even I’m so stupid as to try to wear two pairs of pants, and thanks to years of training I never go outside without wearing pants anymore. If I were to describe this situation using the language of probability theory, I would refer to each pair of pants (i.e., each X) as an **elementary event**. The key characteristic of elementary events is that every time we make an observation (e.g., every time I put on a pair of pants), then the outcome will be one and only one of these events. Like I said, these days I always wear exactly one pair of pants, so my pants satisfy this constraint. Similarly, the set of all possible events is called a **sample space**. Granted, some people would call it a “wardrobe”, but that’s because they’re refusing to think about my pants in probabilistic terms. Sad.

Okay, now that we have a sample space (a wardrobe), which is built from lots of possible elementary events (pants), what we want to do is assign a **probability** of one of these elementary events. For an event X , the probability of that event $P(X)$ is a number that lies between 0 and 1. The bigger the value of $P(X)$, the more likely the event is to occur. So, for example, if $P(X) = 0$, it means the event X is impossible (i.e., I never wear those pants). On the other hand, if $P(X) = 1$ it means that event X is certain to occur (i.e., I always wear those pants). For probability values in the middle, it means that I sometimes wear those pants. For instance, if $P(X) = 0.5$ it means that I wear those pants half of the time.

At this point, we’re almost done. The last thing we need to recognise is that “something always happens”. Every time I put on pants, I really do end up wearing pants (crazy, right?). What this somewhat trite statement means, in probabilistic terms, is that the probabilities of the elementary events all need to add up to 1: which is known as the **law of total probability**, not that any of us really care. More importantly, if these requirements are satisfied, then what we have is a **probability distribution**. For example, this is an example of a probability distribution

Which pants?	Label	Probability
Blue jeans	X_1	$P(X_1) = .5$
Grey jeans	X_2	$P(X_2) = .3$
Black jeans	X_3	$P(X_3) = .1$
Black suit	X_4	$P(X_4) = 0$
Blue tracksuit	X_5	$P(X_5) = .1$

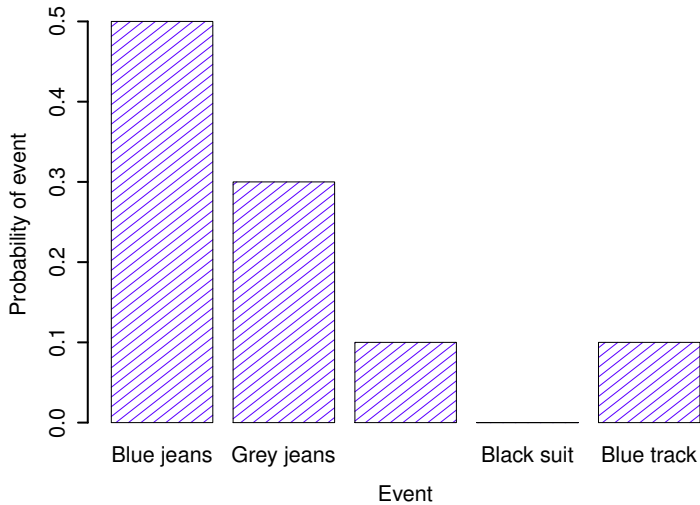


Figure 9.1: A visual depiction of the “pants” probability distribution. There are five “elementary events”, corresponding to the five pairs of pants that I own. Each event has some probability of occurring: this probability is a number between 0 to 1. The sum of these probabilities is 1.

Each of the events has a probability that lies between 0 and 1, and if we add up the probability of all events, they sum to 1. Awesome. We can even draw a nice bar graph (see Section 6.7) to visualise this distribution, as shown in Figure 9.1. And at this point, we’ve all achieved something. You’ve learned what a probability distribution is, and I’ve finally managed to find a way to create a graph that focuses entirely on my pants. Everyone wins!

The only other thing that I need to point out is that probability theory allows you to talk about **non elementary events** as well as elementary ones. The easiest way to illustrate the concept is with an example. In the pants example, it’s perfectly legitimate to refer to the probability that I wear jeans. In this scenario, the “Dan wears jeans” event said to have happened as long as the elementary event that actually did occur is one of the appropriate ones; in this case “blue jeans”, “black jeans” or “grey jeans”. In mathematical terms, we defined the “jeans” event E to correspond to the set of elementary events (X_1, X_2, X_3) . If any of these elementary events occurs, then E is also said to have occurred. Having decided to write down the definition of the E this way, it’s pretty straightforward to state what the probability $P(E)$ is: we just add everything up. In this particular case

$$P(E) = P(X_1) + P(X_2) + P(X_3)$$

and, since the probabilities of blue, grey and black jeans respectively are .5, .3 and .1, the probability that I wear jeans is equal to .9.

At this point you might be thinking that this is all terribly obvious and simple and in a lot of ways that’s true. All we’ve really done is try to wrap some basic mathematics around a few everyday common sense intuitions. However, from these very simple beginnings it’s possible to construct some extremely powerful mathematical tools. I’m definitely not going to go into the details in this book, but what I will do is list – in Table 9.1 – some of the other rules that probabilities satisfy. These rules can be derived from the simple assumptions that I’ve outlined above, but since we don’t actually use these rules for anything in this book, I won’t do so here.

Table 9.1: Some basic rules that probabilities must satisfy. You don't really need to know these rules in order to understand the analyses that we'll talk about later in the book, but they are important if you want to understand probability theory a bit more deeply.

English	Notation	Formula
not A	$P(\neg A)$	$= 1 - P(A)$
A or B	$P(A \cup B)$	$= P(A) + P(B) - P(A \cap B)$
A and B	$P(A \cap B)$	$= P(A B)P(B)$

9.3

The binomial distribution

As you might imagine, probability distributions vary enormously, and there's an enormous range of distributions out there. However, they aren't all equally important. In fact, the vast majority of the content in this book relies on one of five distributions: the binomial distribution, the normal distribution, the t distribution, the χ^2 ("chi-square") distribution and the F distribution. Given this, what I'll do over the next few sections is provide a brief introduction to all five of these, paying special attention to the binomial and the normal. I'll start with the binomial distribution, since it's the simplest of the five.

9.3.1 Introducing the binomial

The theory of probability originated in the attempt to describe how games of chance work, so it seems fitting that our discussion of the **binomial distribution** should involve a discussion of rolling dice and flipping coins. Let's imagine a simple "experiment": in my hot little hand I'm holding 20 identical six-sided dice. On one face of each die there's a picture of a skull; the other five faces are all blank. If I proceed to roll all 20 dice, what's the probability that I'll get exactly 4 skulls? Assuming that the dice are fair, we know that the chance of any one die coming up skulls is 1 in 6; to say this another way, the skull probability for a single die is approximately .167. This is enough information to answer our question, so let's have a look at how it's done.

As usual, we'll want to introduce some names and some notation. We'll let N denote the number of dice rolls in our experiment; which is often referred to as the **size parameter** of our binomial distribution. We'll also use θ to refer to the the probability that a single die comes up skulls, a quantity that is usually called the **success probability** of the binomial.¹ Finally, we'll use X to refer to the results of our experiment, namely the number of skulls I get when I roll the dice. Since the actual value of X is due to chance, we refer to it as a **random variable**. In any case, now that we have all this terminology and notation, we can use it to state the problem a little more precisely. The quantity that we want to calculate is the probability that $X = 4$ given that we know that $\theta = .167$ and $N = 20$. The general "form" of the thing I'm interested in calculating could be written as

$$P(X | \theta, N)$$

and we're interested in the special case where $X = 4$, $\theta = .167$ and $N = 20$. There's only one more

¹Note that the term "success" is pretty arbitrary, and doesn't actually imply that the outcome is something to be desired. If θ referred to the probability that any one passenger gets injured in a bus crash, I'd still call it the success probability, but that doesn't mean I want people to get hurt in bus crashes!

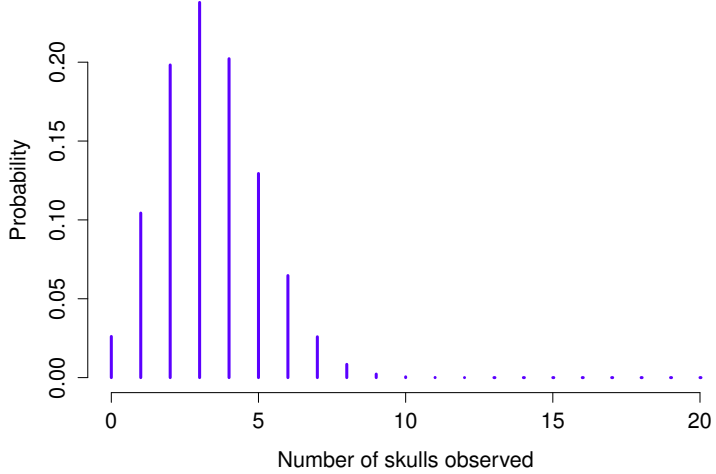


Figure 9.2: The binomial distribution with size parameter of $N = 20$ and an underlying success probability of $\theta = 1/6$. Each vertical bar depicts the probability of one specific outcome (i.e., one possible value of X). Because this is a probability distribution, each of the probabilities must be a number between 0 and 1, and the heights of the bars must sum to 1 as well.

piece of notation I want to refer to before moving on to discuss the solution to the problem. If I want to say that X is generated randomly from a binomial distribution with parameters θ and N , the notation I would use is as follows:

$$X \sim \text{Binomial}(\theta, N)$$

Yeah, yeah. I know what you're thinking: notation, notation, notation. Really, who cares? Very few readers of this book are here for the notation, so I should probably move on and talk about how to use the binomial distribution. I've included the formula for the binomial distribution in Table 9.2, since some readers may want to play with it themselves, but since most people probably don't care that much and because we don't need the formula in this book, I won't talk about it in any detail. Instead, I just want to show you what the binomial distribution looks like. To that end, Figure 9.2 plots the binomial probabilities for all possible values of X for our dice rolling experiment, from $X = 0$ (no skulls) all the way up to $X = 20$ (all skulls). Note that this is basically a bar chart, and is no different to the "pants probability" plot I drew in Figure 9.1. On the horizontal axis we have all the possible events, and on the vertical axis we can read off the probability of each of those events. So, the probability of rolling 4 skulls out of 20 times is about 0.20 (the actual answer is 0.2022036, as we'll see in a moment). In other words, you'd expect that to happen about 20% of the times you repeated this experiment.

9.3.2 Working with the binomial distribution in R

Although some people find it handy to know the formulas in Table 9.2, most people just want to know how to use the distributions without worrying too much about the maths. To that end, R has a function called `dbinom()` that calculates binomial probabilities for us. The main arguments to the function are

Table 9.2: Formulas for the binomial and normal distributions. We don't really use these formulas for anything in this book, but they're pretty important for more advanced work, so I thought it might be best to put them here in a table, where they can't get in the way of the text. In the equation for the binomial, $X!$ is the factorial function (i.e., multiply all whole numbers from 1 to X), and for the normal distribution “exp” refers to the exponential function, which we discussed in Chapter 7. If these equations don't make a lot of sense to you, don't worry too much about them.

$$\begin{array}{ll} \text{Binomial} & \text{Normal} \\ P(X | \theta, N) = \frac{N!}{X!(N-X)!} \theta^X (1-\theta)^{N-X} & p(X | \mu, \sigma) = \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(X-\mu)^2}{2\sigma^2}\right) \end{array}$$

Table 9.3: The naming system for R probability distribution functions. Every probability distribution implemented in R is actually associated with four separate functions, and there is a pretty standardised way for naming these functions.

what it does	prefix	normal distribution	binomial distribution
probability (density) of	d	dnorm()	dbinom()
cumulative probability of	p	pnorm()	pbinom()
generate random number from	r	rnorm()	rbinom()
quantile of	q	qnorm()	qbinom()

- **x**. This is a number, or vector of numbers, specifying the outcomes whose probability you're trying to calculate.
- **size**. This is a number telling R the size of the experiment.
- **prob**. This is the success probability for any one trial in the experiment.

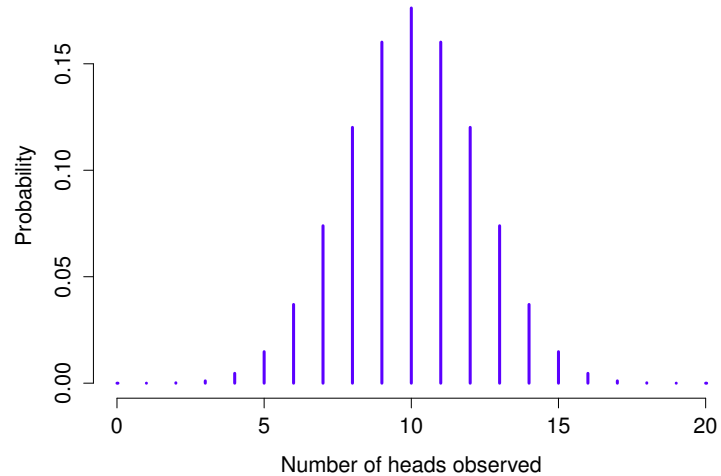
So, in order to calculate the probability of getting `x = 4` skulls, from an experiment of `size = 20` trials, in which the probability of getting a skull on any one trial is `prob = 1/6` ... well, the command I would use is simply this:

```
> dbinom( x = 4, size = 20, prob = 1/6 )
[1] 0.2022036
```

To give you a feel for how the binomial distribution changes when we alter the values of θ and N , let's suppose that instead of rolling dice, I'm actually flipping coins. This time around, my experiment involves flipping a fair coin repeatedly, and the outcome that I'm interested in is the number of heads that I observe. In this scenario, the success probability is now $\theta = 1/2$. Suppose I were to flip the coin $N = 20$ times. In this example, I've changed the success probability, but kept the size of the experiment the same. What does this do to our binomial distribution? Well, as Figure 9.3a shows, the main effect of this is to shift the whole distribution, as you'd expect. Okay, what if we flipped a coin $N = 100$ times? Well, in that case, we get Figure 9.3b. The distribution stays roughly in the middle, but there's a bit more variability in the possible outcomes.

At this point, I should probably explain the name of the `dbinom()` function. Obviously, the “binom” part comes from the fact that we're working with the binomial distribution, but the “d” prefix is probably a bit of a mystery. In this section I'll give a partial explanation: specifically, I'll explain why there is a prefix. As for why it's a “d” specifically, you'll have to wait until the next section. What's going on here

(a)



(b)

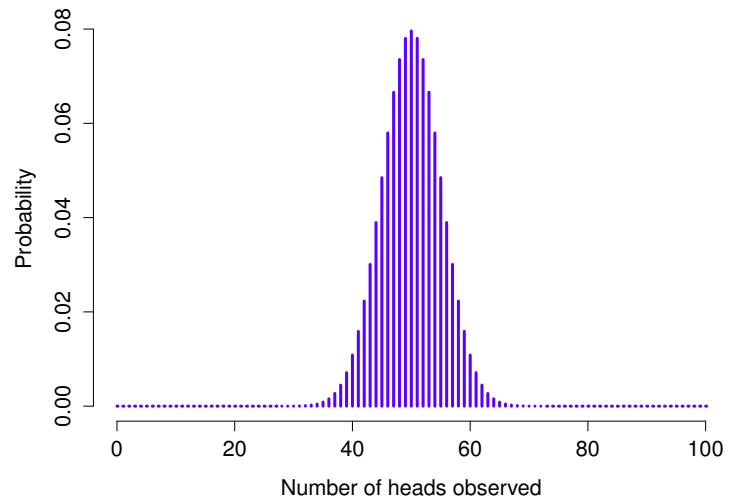


Figure 9.3: Two binomial distributions, involving a scenario in which I'm flipping a fair coin, so the underlying success probability is $\theta = 1/2$. In panel (a), we assume I'm flipping the coin $N = 20$ times. In panel (b) we assume that the coin is flipped $N = 100$ times.

.....

is that R actually provides *four* functions in relation to the binomial distribution. These four functions are `dbinom()`, `pbinom()`, `rbinom()` and `qbinom()`, and each one calculates a different quantity of interest. Not only that, R does the same thing for *every* probability distribution that it implements. No matter what distribution you're talking about, there's a `d` function, a `p` function, a `q` function and a `r` function. This is illustrated in Table 9.3, using the binomial distribution and the normal distribution as examples.

Let's have a look at what all four functions do. Firstly, all four versions of the function require you to specify the `size` and `prob` arguments: no matter what you're trying to get R to calculate, it needs to know what the parameters are. However, they differ in terms of what the other argument is, and what the output is. So let's look at them one at a time.

- The `d` form we've already seen: you specify a particular outcome `x`, and the output is the probability of obtaining exactly that outcome. (the “d” is short for **density**, but ignore that for now).
- The `p` form calculates the **cumulative probability**. You specify a particular quantile `q`, and it tells you the probability of obtaining an outcome *smaller than or equal to* `q`.
- The `q` form calculates the **quantiles** of the distribution. You specify a probability value `p`, and gives you the corresponding percentile. That is, the value of the variable for which there's a probability `p` of obtaining an outcome lower than that value.
- The `r` form is a **random number generator**: specifically, it generates `n` random outcomes from the distribution.

This probably seems a little abstract, so let's look at some concrete examples. Again, we've already covered `dbinom()` so let's focus on the other three versions. We'll start with `pbinom()`, and we'll go back to the skull-dice example. Again, I'm rolling 20 dice, and each die has a 1 in 6 chance of coming up skulls. Suppose, however, that I want to know the probability of rolling 4 *or fewer* skulls. If I wanted to, I could use the `dbinom()` function to calculate the exact probability of rolling 0 skulls, 1 skull, 2 skulls, 3 skulls and 4 skulls and then add these up, but there's a faster way. Instead, I can calculate this using the `pbinom()` function. Here's the command:

```
> pbinom( q= 4, size = 20, prob = 1/6)
[1] 0.7687492
```

In other words, there is a 76.9% chance that I will roll 4 or fewer skulls. Or, to put it another way, R is telling us that a value of 4 is actually the 76.9th percentile of this binomial distribution.

Next, let's consider the `qbinom()` function. Let's say I want to calculate the 75th percentile of the binomial distribution. If we're sticking with our skulls example, I would use the following command to do this:

```
> qbinom( p = 0.75, size = 20, prob = 1/6 )
[1] 4
```

Hm. There's something odd going on here. Let's think this through. What the `qbinom()` function appears to be telling us is that the 75th percentile of the binomial distribution is 4, even though we saw from the `pbinom()` function that 4 is *actually* the 76.9th percentile. And it's definitely the `pbinom()` function that is correct. I promise. The weirdness here comes from the fact that our binomial distribution doesn't really *have* a 75th percentile. Not really. Why not? Well, there's a 56.7% chance of rolling 3 or fewer skulls (you can type `pbinom(3, 20, 1/6)` to confirm this if you want), and a 76.9% chance of rolling 4 or fewer skulls. So there's a sense in which the 75th percentile should lie “in between” 3 and 4 skulls. But that makes no sense at all! You can't roll 20 dice and get 3.9 of them come up skulls. This issue can be handled in different ways: you could report an in between value (or *interpolated* value, to use

the technical name) like 3.9, you could round down (to 3) or you could round up (to 4). The `qbinom()` function rounds upwards: if you ask for a percentile that doesn't actually exist (like the 75th in this example), R finds the smallest value for which the percentile rank is *at least* what you asked for. In this case, since the “true” 75th percentile (whatever that would mean) lies somewhere between 3 and 4 skulls, R rounds up and gives you an answer of 4. This subtlety is tedious, I admit, but thankfully it's only an issue for discrete distributions like the binomial (see Section 2.2.5 for a discussion of continuous versus discrete). The other distributions that I'll talk about (normal, t , χ^2 and F) are all continuous, and so R can always return an exact quantile whenever you ask for it.

Finally, we have the random number generator. To use the `rbinom()` function, you specify how many times R should “simulate” the experiment using the `n` argument, and it will generate random outcomes from the binomial distribution. So, for instance, suppose I were to repeat my die rolling experiment 100 times. I could get R to simulate the results of these experiments by using the following command:

```
> rbinom( n = 100, size = 20, prob = 1/6 )
[1] 3 4 8 4 4 3 1 4 4 1 3 0 3 3 4 3 1 2 2 3 4 2
[23] 2 1 2 4 4 3 5 4 2 3 1 2 7 4 2 5 2 3 0 2 3 3
[45] 2 2 2 3 4 4 2 0 2 4 4 3 4 1 2 3 5 3 7 5 0 5
[67] 1 5 4 3 4 4 1 5 4 4 3 2 3 3 4 5 0 5 1 4 7 2
[89] 5 1 1 2 4 5 5 3 3 3 3 3
```

As you can see, these numbers are pretty much what you'd expect given the distribution shown in Figure 9.2. Most of the time I roll somewhere between 1 to 5 skulls. There are a lot of subtleties associated with random number generation using a computer,² but for the purposes of this book we don't need to worry too much about them.

9.4

The normal distribution

While the binomial distribution is conceptually the simplest distribution to understand, it's not the most important one. That particular honour goes to the **normal distribution**, which is also referred to as “the bell curve” or a “Gaussian distribution”. A normal distribution is described using two parameters, the mean of the distribution μ and the standard deviation of the distribution σ . The notation that we sometimes use to say that a variable X is normally distributed is as follows:

$$X \sim \text{Normal}(\mu, \sigma)$$

Of course, that's just notation. It doesn't tell us anything interesting about the normal distribution itself. As was the case with the binomial distribution, I have included the formula for the normal distribution in this book, because I think it's important enough that everyone who learns statistics should at least look at it, but since this is an introductory text I don't want to focus on it, so I've tucked it away in Table 9.2. Similarly, the R functions for the normal distribution are `dnorm()`, `pnorm()`, `qnorm()` and `rnorm()`. However, they behave in pretty much exactly the same way as the corresponding functions for the binomial distribution, so there's not a lot that you need to know. The only thing that I should point out is that the argument names for the parameters are `mean` and `sd`. In pretty much every other respect, there's nothing else to add.

²Since computers are deterministic machines, they can't actually produce truly random behaviour. Instead, what they do is take advantage of various mathematical functions that share a lot of similarities with true randomness. What this means is that any random numbers generated on a computer are *pseudorandom*, and the quality of those numbers depends on the specific method used. By default R uses the “Mersenne twister” method. In any case, you can find out more by typing `?Random`, but as usual the R help files are fairly dense.

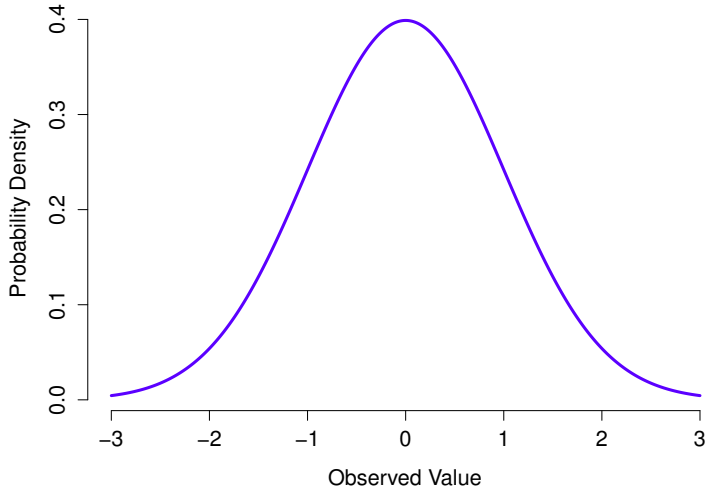


Figure 9.4: The normal distribution with mean $\mu = 0$ and standard deviation $\sigma = 1$. The x -axis corresponds to the value of some variable, and the y -axis tells us something about how likely we are to observe that value. However, notice that the y -axis is labelled “Probability Density” and not “Probability”. There is a subtle and somewhat frustrating characteristic of continuous distributions that makes the y axis behave a bit oddly: the height of the curve here isn’t actually the probability of observing a particular x value. On the other hand, it *is* true that the heights of the curve tells you which x values are more likely (the higher ones!). (see Section 9.4.1 for all the annoying details)

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Instead of focusing on the maths, let’s try to get a sense for what it means for a variable to be normally distributed. To that end, have a look at Figure 9.4, which plots a normal distribution with mean $\mu = 0$ and standard deviation $\sigma = 1$. You can see where the name “bell curve” comes from: it looks a bit like a bell. Notice that, unlike the plots that I drew to illustrate the binomial distribution, the picture of the normal distribution in Figure 9.4 shows a smooth curve instead of “histogram-like” bars. This isn’t an arbitrary choice: the normal distribution is continuous, whereas the binomial is discrete. For instance, in the die rolling example from the last section, it was possible to get 3 skulls or 4 skulls, but impossible to get 3.9 skulls. The figures that I drew in the previous section reflected this fact: in Figure 9.2, for instance, there’s a bar located at $X = 3$ and another one at $X = 4$, but there’s nothing in between. Continuous quantities don’t have this constraint. For instance, suppose we’re talking about the weather. The temperature on a pleasant Spring day could be 23 degrees, 24 degrees, 23.9 degrees, or anything in between since temperature is a continuous variable, and so a normal distribution might be quite appropriate for describing Spring temperatures.³

With this in mind, let’s see if we can’t get an intuition for how the normal distribution works. Firstly, let’s have a look at what happens when we play around with the parameters of the distribution. To that end, Figure 9.5 plots normal distributions that have different means, but have the same standard deviation. As you might expect, all of these distributions have the same “width”. The only difference between them is that they’ve been shifted to the left or to the right. In every other respect they’re

³In practice, the normal distribution is so handy that people tend to use it even when the variable isn’t actually continuous. As long as there are enough categories (e.g., Likert scale responses to a questionnaire), it’s pretty standard practice to use the normal distribution as an approximation. This works out much better in practice than you’d think.

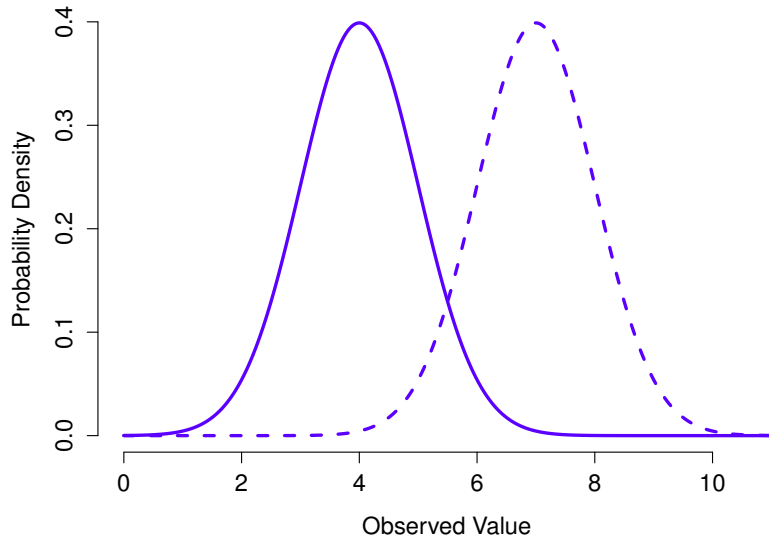


Figure 9.5: An illustration of what happens when you change the mean of a normal distribution. The solid line depicts a normal distribution with a mean of $\mu = 4$. The dashed line shows a normal distribution with a mean of $\mu = 7$. In both cases, the standard deviation is $\sigma = 1$. Not surprisingly, the two distributions have the same shape, but the dashed line is shifted to the right.

identical. In contrast, if we increase the standard deviation while keeping the mean constant, the peak of the distribution stays in the same place, but the distribution gets wider, as you can see in Figure 9.6. Notice, though, that when we widen the distribution, the height of the peak shrinks. This has to happen: in the same way that the heights of the bars that we used to draw a discrete binomial distribution have to *sum* to 1, the total *area under the curve* for the normal distribution must equal 1. Before moving on, I want to point out one important characteristic of the normal distribution. Irrespective of what the actual mean and standard deviation are, 68.3% of the area falls within 1 standard deviation of the mean. Similarly, 95.4% of the distribution falls within 2 standard deviations of the mean, and 99.7% of the distribution is within 3 standard deviations. This idea is illustrated in Figure 9.7.

9.4.1 Probability density

There's something I've been trying to hide throughout my discussion of the normal distribution, something that some introductory textbooks omit completely. They might be right to do so: this "thing" that I'm hiding is a bit weird and a little counterintuitive, even by the admittedly distorted standards that apply in statistics. With that in mind, I should warn you that this section is a little hard to follow. Fortunately, it's not something that you need to understand at a deep level in order to do basic statistics: rather, it's something that starts to become important later on when you move beyond the basics. So, if it doesn't make complete sense, don't worry: try to make sure that you follow the gist of it.

Throughout my discussion of the normal distribution, there's been one or two things that don't quite make sense. Perhaps you noticed that the y -axis in these figures is labelled "Probability Density" rather

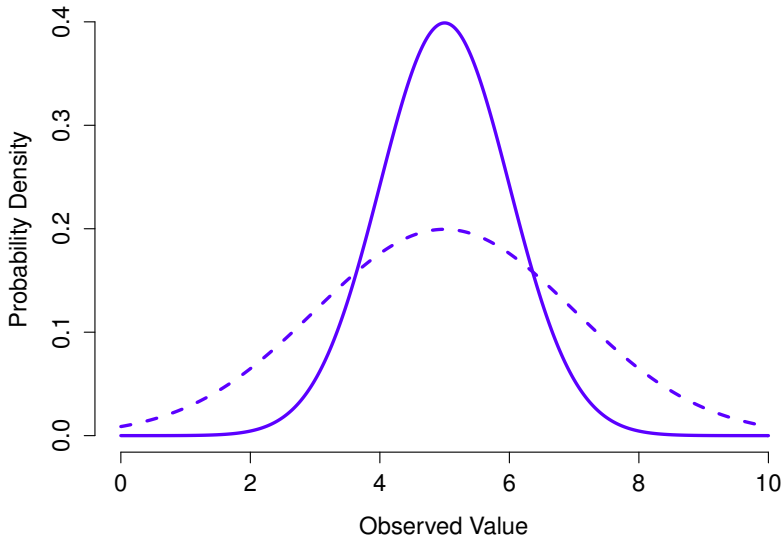


Figure 9.6: An illustration of what happens when you change the standard deviation of a normal distribution. Both distributions plotted in this figure have a mean of $\mu = 5$, but they have different standard deviations. The solid line plots a distribution with standard deviation $\sigma = 1$, and the dashed line shows a distribution with standard deviation $\sigma = 2$. As a consequence, both distributions are “centred” on the same spot, but the dashed line is wider than the solid one.

than density. Maybe you noticed that I used $p(X)$ instead of $P(X)$ when giving the formula for the normal distribution. Maybe you’re wondering why R uses the “d” prefix for functions like `dnorm()`. And maybe, just maybe, you’ve been playing around with the `dnorm()` function, and you accidentally typed in a command like this:

```
> dnorm( x = 1, mean = 1, sd = 0.1 )
[1] 3.989423
```

And if you’ve done the last part, you’re probably very confused. I’ve asked R to calculate the probability that `x = 1`, for a normally distributed variable with `mean = 1` and standard deviation `sd = 0.1`; and it tells me that the probability is 3.99. But, as we discussed earlier, probabilities *can’t* be larger than 1. So either I’ve made a mistake, or that’s not a probability.

As it turns out, the second answer is correct. What we’ve calculated here isn’t actually a probability: it’s something else. To understand what that something is, you have to spend a little time thinking about what it really *means* to say that X is a continuous variable. Let’s say we’re talking about the temperature outside. The thermometer tells me it’s 23 degrees, but I know that’s not really true. It’s not *exactly* 23 degrees. Maybe it’s 23.1 degrees, I think to myself. But I know that that’s not really true either, because it might actually be 23.09 degrees. But, I know that... well, you get the idea. The tricky thing with genuinely continuous quantities is that you never really know exactly what they are.

Now think about what this implies when we talk about probabilities. Suppose that tomorrow’s maximum temperature is sampled from a normal distribution with mean 23 and standard deviation 1. What’s the probability that the temperature will be *exactly* 23 degrees? The answer is “zero”, or

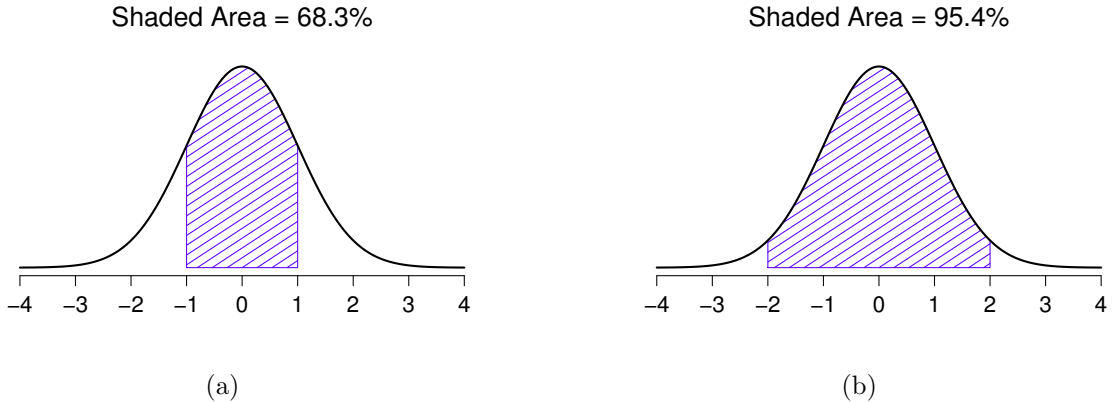


Figure 9.7: The area under the curve tells you the probability that an observation falls within a particular range. The solid lines plot normal distributions with mean $\mu = 0$ and standard deviation $\sigma = 1$. The shaded areas illustrate “areas under the curve” for two important cases. In panel a, we can see that there is a 68.3% chance that an observation will fall within one standard deviation of the mean. In panel b, we see that there is a 95.4% chance that an observation will fall within two standard deviations of the mean.

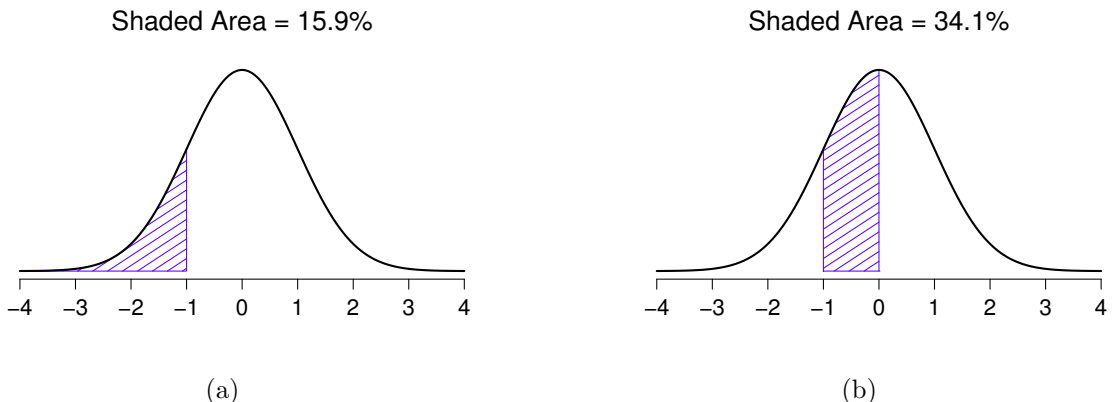


Figure 9.8: Two more examples of the “area under the curve idea”. There is a 15.9% chance that an observation is one standard deviation below the mean or smaller (panel a), and a 34.1% chance that the observation is greater than one standard deviation below the mean but still below the mean (panel b). Notice that if you add these two numbers together you get $15.9\% + 34.1\% = 50\%$. For normally distributed data, there is a 50% chance that an observation falls below the mean. And of course that also implies that there is a 50% chance that it falls above the mean.

possibly, “a number so close to zero that it might as well be zero”. Why is this? It’s like trying to throw a dart at an infinitely small dart board: no matter how good your aim, you’ll never hit it. Similarly, you’ll never get a value of exactly 23. It’ll always be something like 23.1 or 22.99998 or something. In other words, it’s completely meaningless to talk about the probability that the temperature is exactly 23 degrees. However, in everyday language, if I told you that it was 23 degrees outside and it turned out to be 22.9998 degrees, you probably wouldn’t call me a liar. Because in everyday language, “23 degrees” really means something like “somewhere between 22.5 and 23.5 degrees”. And while it doesn’t feel very meaningful to ask about the probability that the temperature is exactly 23 degrees, it does seem sensible to ask about the probability that the temperature lies between 22.5 and 23.5, or between 20 and 30, or any other range of temperatures.

The point of this discussion is to make clear that, when we’re talking about continuous distributions, it’s not meaningful to talk about the probability of a specific value. However, what we *can* talk about is the probability that the value lies within a particular range of values. To find out the probability associated with a particular range, what you need to do is calculate the “area under the curve”. We’ve seen this concept already: in Figure 9.7, the shaded areas shown depict genuine probabilities (e.g., in Figure 9.7a it shows the probability of observing a value that falls within 1 standard deviation of the mean).

Okay, so that explains part of the story. I’ve explained a little bit about how continuous probability distributions should be interpreted (i.e., area under the curve is the key thing), but I haven’t actually explained what the `dnorm()` function actually calculates. Equivalently, what does the formula for $p(x)$ that I described earlier actually mean? Obviously, $p(x)$ doesn’t describe a probability, but what is it? The name for this quantity $p(x)$ is a **probability density**, and in terms of the plots we’ve been drawing, it corresponds to the *height* of the curve. The densities themselves aren’t meaningful in and of themselves: but they’re “rigged” to ensure that the *area* under the curve is always interpretable as genuine probabilities. To be honest, that’s about as much as you really need to know for now.⁴

9.5

Other useful distributions

The normal distribution is the distribution that statistics makes most use of (for reasons to be discussed shortly), and the binomial distribution is a very useful one for lots of purposes. But the world of statistics is filled with probability distributions, some of which we’ll run into in passing. In particular, the three that will appear in this book are the t distribution, the χ^2 distribution and the F distribution. I won’t give formulas for any of these, or talk about them in too much detail, but I will show you some pictures.

- The **t distribution** is a continuous distribution that looks very similar to a normal distribution, but has heavier tails: see Figure 9.9. This distribution tends to arise in situations where you think that the data actually follow a normal distribution, but you don’t know the mean or standard deviation. As you might expect, the relevant R functions are `dt()`, `pt()`, `qt()` and `rt()`, and we’ll run into this distribution again in Chapter 13.
- The **χ^2 distribution** is another distribution that turns up in lots of different places. The situation in which we’ll see it is when doing categorical data analysis (Chapter 12), but it’s one of those things that actually pops up all over the place. When you dig into the maths (and who doesn’t love

⁴For those readers who know a little calculus, I’ll give a slightly more precise explanation. In the same way that probabilities are non-negative numbers that must sum to 1, probability densities are non-negative numbers that must integrate to 1 (where the integral is taken across all possible values of X). To calculate the probability that X falls between a and b we calculate the definite integral of the density function over the corresponding range, $\int_a^b p(x) dx$. If you don’t remember or never learned calculus, don’t worry about this. It’s not needed for this book.

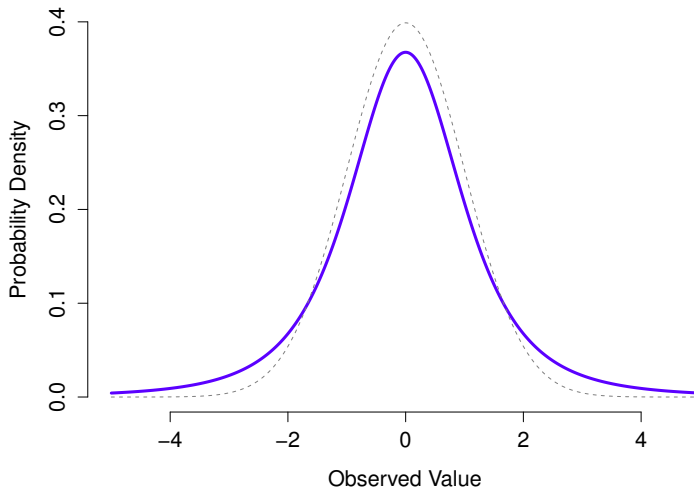


Figure 9.9: A t distribution with 3 degrees of freedom (solid line). It looks similar to a normal distribution, but it's not quite the same. For comparison purposes, I've plotted a standard normal distribution as the dashed line. Note that the “tails” of the t distribution are “heavier” (i.e., extend further outwards) than the tails of the normal distribution? That's the important difference between the two.

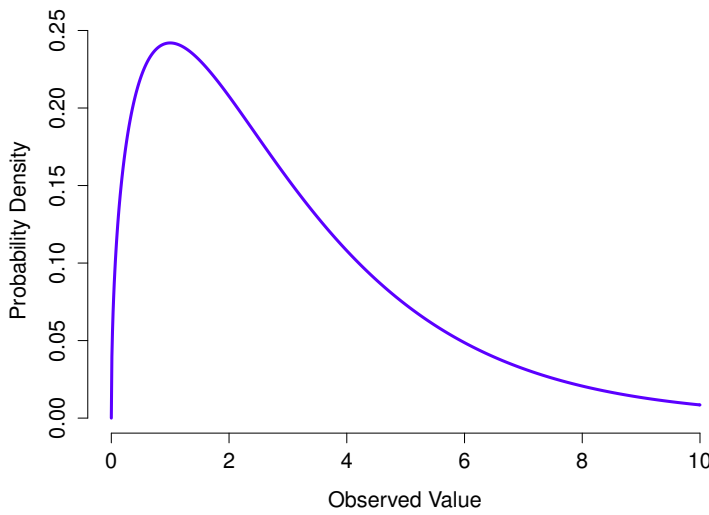


Figure 9.10: A χ^2 distribution with 3 degrees of freedom. Notice that the observed values must always be greater than zero, and that the distribution is pretty skewed. These are the key features of a chi-square distribution.

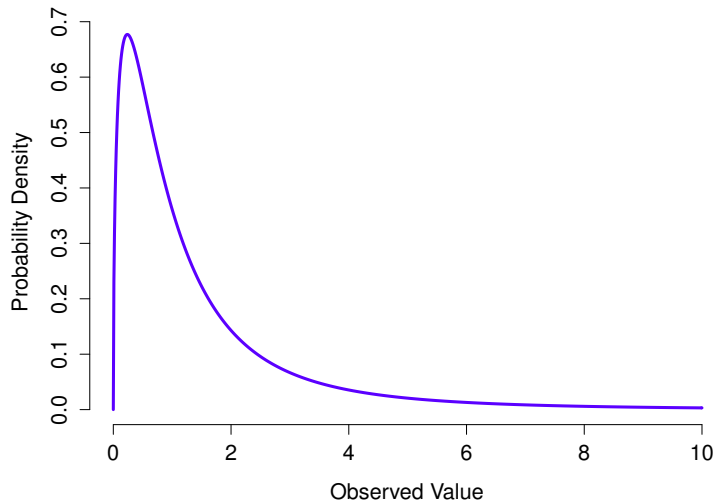


Figure 9.11: An F distribution with 3 and 5 degrees of freedom. Qualitatively speaking, it looks pretty similar to a chi-square distribution, but they're not quite the same in general.

(doing that?), it turns out that the main reason why the χ^2 distribution turns up all over the place is that, if you have a bunch of variables that are normally distributed, square their values and then add them up (a procedure referred to as taking a “sum of squares”), this sum has a χ^2 distribution. You’d be amazed how often this fact turns out to be useful. Anyway, here’s what a χ^2 distribution looks like: Figure 9.10. Once again, the R commands for this one are pretty predictable: `dchisq()`, `pchisq()`, `qchisq()`, `rchisq()`.

- The **F distribution** looks a bit like a χ^2 distribution, and it arises whenever you need to compare two χ^2 distributions to one another. Admittedly, this doesn’t exactly sound like something that any sane person would want to do, but it turns out to be very important in real world data analysis. Remember when I said that χ^2 turns out to be the key distribution when we’re taking a “sum of squares”? Well, what that means is if you want to compare two different “sums of squares”, you’re probably talking about something that has an F distribution. Of course, as yet I still haven’t given you an example of anything that involves a sum of squares, but I will... in Chapter 14. And that’s where we’ll run into the F distribution. Oh, and here’s a picture: Figure 9.11. And of course we can get R to do things with F distributions just by using the commands `df()`, `pf()`, `qf()` and `rf()`.

Because these distributions are all tightly related to the normal distribution and to each other, and because they are will turn out to be the important distributions when doing inferential statistics later in this book, I think it’s useful to do a little demonstration using R, just to “convince ourselves” that these distributions really are related to each other in the way that they’re supposed to be. First, we’ll use the `rnorm()` function to generate 1000 normally-distributed observations:

```
> normal.a <- rnorm( n=1000, mean=0, sd=1 )
> print(normal.a)
[1]  0.2913131706 -0.4156161554  0.1482611948  0.8516858463 -0.6658081840
[6]  0.8827940964  1.3757851963  0.2497812249 -0.1926513775  0.2160192605
```

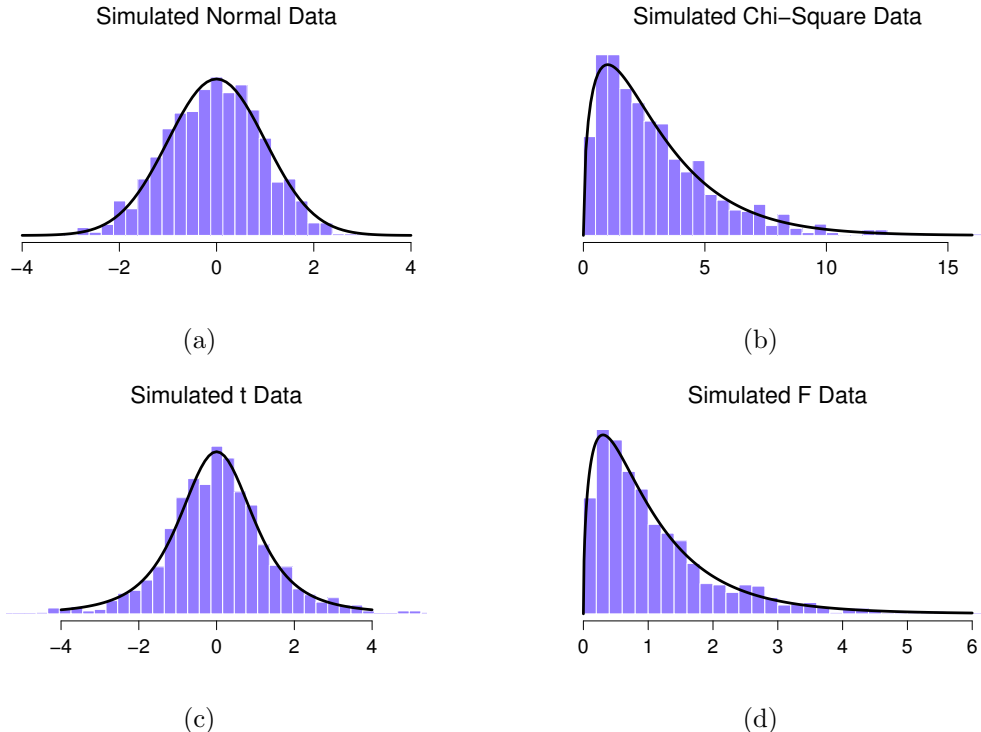


Figure 9.12: Data sampled from different distributions. See the main text for details.

```
[11] -0.7982884040 -1.4027212056 0.0281455244 -0.1266362460 0.8645205990
BLAH BLAH BLAH
```

So the `normal.a` variable contains 1000 numbers that are normally distributed, and have mean 0 and standard deviation 1, and the actual print out of these numbers goes on for rather a long time. Note that, because the default parameters of the `rnorm()` function are `mean=0` and `sd=1`, I could have shortened the command to `rnorm(n=1000)`. In any case, what we can do is use the `hist()` function to draw a histogram of the data, like so:

```
> hist( normal.a )
```

If you do this, you should see something similar to Figure 9.12a. Your plot won't look quite as pretty as the one in the figure, of course, because I've played around with all the formatting (see Chapter 6), and I've also plotted the true distribution of the data as a solid black line (i.e., a normal distribution with mean 0 and standard deviation 1) so that you can compare the data that we just generated to the true distribution.

In the previous example all I did was generate lots of normally distributed observations using `rnorm()` and then compared those to the true probability distribution in the figure (using `dnorm()` to generate the black line in the figure, but I didn't show the commands for that). Now let's try something trickier. We'll try to generate some observations that follow a chi-square distribution with 3 degrees of freedom, but instead of using `rchisq()`, we'll start with variables that are normally distributed, and see if we can exploit the known relationships between normal and chi-square distributions to do the work. As I

mentioned earlier, a chi-square distribution with k degrees of freedom is what you get when you take k normally-distributed variables (with mean 0 and standard deviation 1), square them, and add them up. Since we want a chi-square distribution with 3 degrees of freedom, we'll need to supplement our `normal.a` data with two more sets of normally-distributed observations, imaginatively named `normal.b` and `normal.c`:

```
> normal.b <- rnorm( n=1000 ) # another set of normally distributed data
> normal.c <- rnorm( n=1000 ) # and another!
```

Now that we've done that, the theory says we should square these and add them together, like this

```
> chi.sq.3 <- (normal.a)^2 + (normal.b)^2 + (normal.c)^2
```

and the resulting `chi.sq.3` variable should contain 1000 observations that follow a chi-square distribution with 3 degrees of freedom. You can use the `hist()` function to have a look at these observations yourself, using a command like this,

```
> hist( chi.sq.3 )
```

and you should obtain a result that looks pretty similar to the plot in Figure 9.12b. Once again, the plot that I've drawn is a little fancier: in addition to the histogram of `chi.sq.3`, I've also plotted a chi-square distribution with 3 degrees of freedom. It's pretty clear that – even though I used `rnorm()` to do all the work rather than `rchisq()` – the observations stored in the `chi.sq.3` variable really do follow a chi-square distribution. Admittedly, this probably doesn't seem all that interesting right now, but later on when we start encountering the chi-square distribution in Chapter 12, it will be useful to understand the fact that these distributions are related to one another.

We can extend this demonstration to the t distribution and the F distribution. Earlier, I implied that the t distribution is related to the normal distribution when the standard deviation is unknown. That's certainly true, and that's the what we'll see later on in Chapter 13, but there's a somewhat more precise relationship between the normal, chi-square and t distributions. Suppose we “scale” our chi-square data by dividing it by the degrees of freedom, like so

```
> scaled.chi.sq.3 <- chi.sq.3 / 3
```

we then take a set of normally distributed variables and divide them by (the square root of) our scaled chi-square variable which had $df = 3$, and the result is a t distribution with 3 degrees of freedom:

```
> normal.d <- rnorm( n=1000 ) # yet another set of normally distributed data
> t.3 <- normal.d / sqrt( scaled.chi.sq.3 ) # divide by square root of scaled chi-square to get t
```

If we plot the histogram of `t.3`, we end up with something that looks very similar to Figure 9.12c. Similarly, we can obtain an F distribution by taking the ratio between two scaled chi-square distributions. Suppose, for instance, we wanted to generate data from an F distribution with 3 and 20 degrees of freedom. We could do this using `df()`, but we could also do the same thing by generating two chi-square variables, one with 3 degrees of freedom, and the other with 20 degrees of freedom. As the example with `chi.sq.3` illustrates, we can actually do this using `rnorm()` if we really want to, but this time I'll take a short cut:

```
> chi.sq.20 <- rchisq( 1000, 20 ) # generate chi square data with df = 20...
> scaled.chi.sq.20 <- chi.sq.20 / 20 # scale the chi square variable...
> F.3.20 <- scaled.chi.sq.3 / scaled.chi.sq.20 # take the ratio of the two chi squares...
> hist( F.3.20 ) # ... and draw a picture
```

The resulting [F.3.20](#) variable does in fact store variables that follow an F distribution with 3 and 20 degrees of freedom. This is illustrated in Figure [9.12d](#), which plots the histogram of the observations stored in [F.3.20](#) against the true F distribution with $df_1 = 3$ and $df_2 = 20$. Again, they match.

Okay, time to wrap this section up. We've seen three new distributions: χ^2 , t and F . They're all continuous distributions, and they're all closely related to the normal distribution. I've talked a little bit about the precise nature of this relationship, and shown you some R commands that illustrate this relationship. The key thing for our purposes, however, is not that you have a deep understanding of all these different distributions, nor that you remember the precise relationships between them. The main thing is that you grasp the basic idea that these distributions are all deeply related to one another, and to the normal distribution. Later on in this book, we're going to run into data that are normally distributed, or at least assumed to be normally distributed. What I want you to understand right now is that, if you make the assumption that your data are normally distributed, you shouldn't be surprised to see χ^2 , t and F distributions popping up all over the place when you start trying to do your data analysis.

9.6

What does probability mean?

Now that we've discussed the a little bit of the mechanics and the mathematics of probability, let's have a go at trying to define probability. Specifically, what does "probability" correspond to in the real world? This turns out to be surprisingly difficult. Even though everyone has some intuitive notion of "the chances of an event", and statisticians all agree about how chance should be mapped into the mathematics, they don't actually agree on how to map it onto the world. In other words, the philosophical, physical and psychological meaning of the word "probability" is in serious dispute. Now, in one sense it doesn't matter: it's the maths that does most of the real grunt work in statistics, and we all agree on the maths. However that doesn't make the semantics unimportant. In fact, it's critical, because it influences the way you *interpret* the statistics, and leads to subtle ideological differences about how you're allowed to *use* the mathematics.

So, what might we mean when we say that "the probability of X is 30%"? How shall we define this, without our definition being circular, and without retreating completely into a mathematical definition that bears no connection to the physical world? In this section, I give a brief (and not at all rigorous) introduction the two main approaches that exist in the literature. These are by no means the only approaches, but they're the two big ones.

9.6.1 The frequentist view

The first of the two major approaches to probability, and the more dominant one in statistics, is referred to as the **frequentist view**, and it defines probability as a **long-run frequency**. Suppose we were to try flipping a fair coin, over and over again. By definition, this is a coin that has $P(H) = 0.5$. What might we observe? One possibility is that the first 20 flips might look like this:

T,H,H,H,H,T,T,H,H,H,T,H,T,T,T,T,H

As it happens, 11 of these 20 coin flips (55%) came up heads. Now suppose that I'd been keeping a running tally of the number of heads (which I'll call N_H) that I've seen, across the first N flips, and calculate the proportion of heads N_H/N every time. Here's what I'd get (I did literally flip coins to produce this!):

number of flips	1	2	3	4	5	6	7	8	9	10
number of heads	0	1	2	3	4	4	4	5	6	7
proportion	.00	.50	.67	.75	.80	.67	.57	.63	.67	.70

number of flips	11	12	13	14	15	16	17	18	19	20
number of heads	8	8	9	10	10	10	10	10	10	11
proportion	.73	.67	.69	.71	.67	.63	.59	.56	.53	.55

Notice that at the start of the sequence, the *proportion* of heads fluctuates wildly, starting at .00 and rising as high as .80. Later on, one gets the impression that it dampens out a bit, with more and more of the values actually being pretty close to the “right” answer of .50. This is the frequentist definition of probability in a nutshell: flip a fair coin over and over again, and as N grows large (approaches infinity, denoted $N \rightarrow \infty$), the proportion of heads will converge to 50%. There are some subtle technicalities that the mathematicians care about, but qualitatively speaking, that’s how the frequentists define probability. Unfortunately, I don’t have an infinite number of coins, or the infinite patience required to flip a coin an infinite number of times. However, I do have a computer, and computers excel at mindless repetitive tasks. So I asked my computer to simulate flipping a coin 1000 times, and then drew a picture of what happens to the proportion N_H/N as N increases. Actually, I did it four times, just to make sure it wasn’t a fluke. The results are shown in Figure 9.13. As you can see, the *proportion of observed heads* eventually stops fluctuating, and settles down; when it does, the number at which it finally settles is the true probability of heads.

The frequentist definition of probability has some desirable characteristics. Firstly, it is objective: the probability of an event is *necessarily* grounded in the world. The only way that probability statements can make sense is if they refer to (a sequence of) events that occur in the physical universe.⁵ Secondly, it is unambiguous: any two people watching the same sequence of events unfold, trying to calculate the probability of an event, must inevitably come up with the same answer. However, it also has undesirable characteristics. Firstly, infinite sequences don’t exist in the physical world. Suppose you picked up a coin from your pocket and started to flip it. Every time it lands, it impacts on the ground. Each impact wears the coin down a bit; eventually, the coin will be destroyed. So, one might ask whether it really makes sense to pretend that an “infinite” sequence of coin flips is even a meaningful concept, or an objective one. We can’t say that an “infinite sequence” of events is a real thing in the physical universe, because the physical universe doesn’t allow infinite anything. More seriously, the frequentist definition has a narrow scope. There are lots of things out there that human beings are happy to assign probability to in everyday language, but cannot (even in theory) be mapped onto a hypothetical sequence of events. For instance, when the meteorologist comes on TV and says, “the probability of rain in Adelaide tomorrow, 1 November 2010, is 60%” we humans are happy to accept this. But it’s not clear how to define this in frequentist terms. There’s only one city of Adelaide, and only 1 November 2010. There’s no infinite sequence of events here, just a once-off thing. Frequentist probability genuinely *forbids* us from making probability statements about a single event. From the frequentist perspective, “it will either rain tomorrow, or it will not; there is no probability”. Now, it should be said that there are some very clever tricks that frequentists can use to get around this. One possibility is that what the meteorologist means is something like this: “There is a category of days for which I predict a 60% chance of rain; if we look only across those days for which I make this prediction, then on 60% of those days it will actually rain”. It’s very weird and counterintuitive to think of it this way, but you do see frequentists do this sometimes. And it *will* come up later in this book (see Section 10.5).

⁵This doesn’t mean that frequentists can’t make hypothetical statements, of course; it’s just that if you want to make a *true* statement about probability, then it has to refer to a real world sequence of observations.

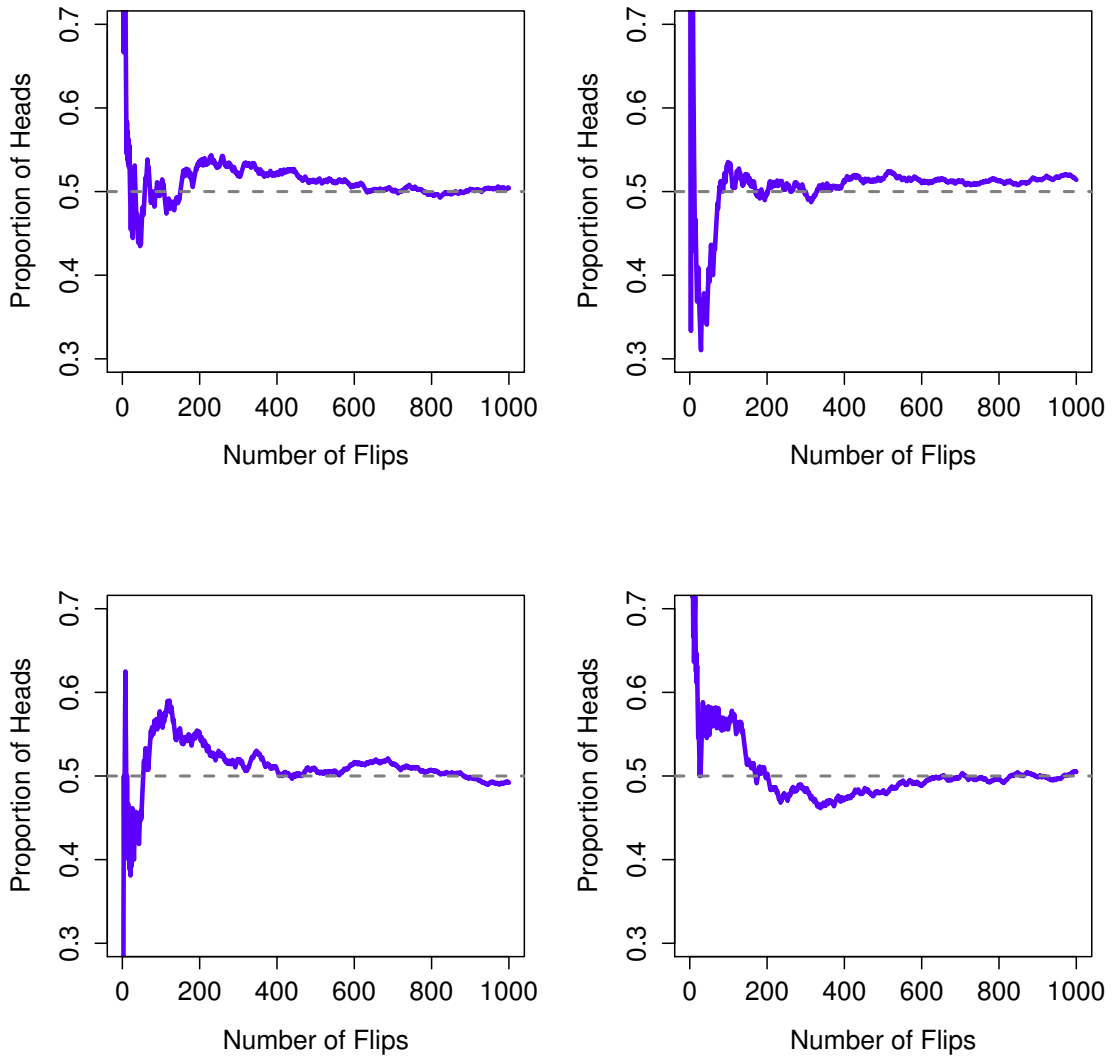


Figure 9.13: An illustration of how frequentist probability works. If you flip a fair coin over and over again, the proportion of heads that you've seen eventually settles down, and converges to the true probability of 0.5. Each panel shows four different simulated experiments: in each case, we pretend we flipped a coin 1000 times, and kept track of the proportion of flips that were heads as we went along. Although none of these sequences actually ended up with an exact value of .5, if we'd extended the experiment for an infinite number of coin flips they would have.

9.6.2 The Bayesian view

The **Bayesian view** of probability is often called the subjectivist view, and it is a minority view among statisticians, but one that has been steadily gaining traction for the last several decades. There are many flavours of Bayesianism, making hard to say exactly what “the” Bayesian view is. The most common way of thinking about subjective probability is to define the probability of an event as the **degree of belief** that an intelligent agent assigns to that truth of that event. From that perspective, probabilities don’t exist in the world, but rather in the thoughts and assumptions of people. However, in order for this approach to work, we need some way of operationalising “degree of belief”. One way that you can do this is to formalise it in terms of “rational gambling”, though there are many other ways. Suppose that I believe that there’s a 60% probability of rain tomorrow. If someone offers me a bet: if it rains tomorrow, then I win \$5, but if it doesn’t rain then I lose \$5. Clearly, from my perspective, this is a pretty good bet. On the other hand, if I think that the probability of rain is only 40%, then it’s a bad bet to take. Thus, we can operationalise the notion of a “subjective probability” in terms of what bets I’m willing to accept.

What are the advantages and disadvantages to the Bayesian approach? The main advantage is that it allows you to assign probabilities to any event you want to. You don’t need to be limited to those events that are repeatable. The main disadvantage (to many people) is that we can’t be purely objective – specifying a probability requires us to specify an entity that has the relevant degree of belief. This entity might be a human, an alien, a robot, or even a statistician, but there has to be an intelligent agent out there that believes in things. To many people this is uncomfortable: it seems to make probability arbitrary. While the Bayesian approach does require that the agent in question be rational (i.e., obey the rules of probability), it does allow everyone to have their own beliefs; I can believe the coin is fair and you don’t have to, even though we’re both rational. The frequentist view doesn’t allow any two observers to assign different probabilities to the same event; the Bayesian view does. And while there are some attempts out there to “fix” this none are quite convincing. In short, where the frequentist view is sometimes considered to be too narrow (forbids lots of things that we want to assign probabilities to), the Bayesian view is often thought to be too broad (allows too many differences between observers). I personally prefer the Bayesian view, but the majority of statistical analyses are based on the frequentist view, so this book doesn’t go into the Bayesian view in any detail.

9.7

Summary

- Probability theory versus statistics (Section 9.1)
- Basics of probability theory (Section 9.2)
- Binomial distribution (Section 9.3), normal distribution (Section 9.4), and others (Section 9.5)
- Frequentist versus Bayesian views of probability (Section 9.6)

10. Estimating population parameters from a sample

Probability theory is a powerful tool: and without probability theory it would be more or less impossible to do inferential statistics. However, there's more to statistical inference than just probability theory. In this chapter we'll move away from talking purely about probabilities, and instead start talking about making statistical inferences. There are two big ideas in inferential statistics: estimation and hypothesis testing. This chapter discusses estimation.

10.1

Samples, populations and sampling

In order to talk about “making inferences” at all, we need to be a bit more explicit about what it is that we’re drawing inferences *from* (the sample) and what it is that we’re drawing inferences *about* (the population).

In almost every situation of interest, what we have available to us as researchers is a **sample** of data. We might have run experiment with some number of participants; a polling company might have phoned some number of people to ask questions about voting intentions; etc. Regardless: the data set available to us is finite, and incomplete. We can’t possibly get every person in the world to do our experiment; a polling company doesn’t have the time or the money to ring up every voter in the country etc. In our earlier discussion of descriptive statistics (Chapter 5), this sample was the only thing we were interested in. Our only goal was to find ways of describing, summarising and graphing that sample. This is about to change.

A sample is a concrete thing. You can open up a data file, and there’s the data from your sample. A **population**, on the other hand, is a more abstract idea. It refers to the set of all possible people, or all possible observations, that you want to draw conclusions about. Almost always, the population is *much* bigger than the sample. For instance, in our “polling company” example previously, the population of interest consists of all enrolled voters – millions of people. In contrast, the sample would probably only consist of 1000 people or so. For a psychological experiment, the situation is more complex. Suppose I run an experiment using 100 undergraduate students as my participants. My goal, as a cognitive scientist, is to try to learn something about how the mind works. So, which of the following would count as “the population”:

- All of the undergraduate psychology students at the University of Adelaide?
- Undergraduate psychology students in general, anywhere in the world?
- Australians currently living?

- Australians of similar ages to my sample?
- Anyone currently alive?
- Any human being, past, present or future?
- Any biological organism with a sufficient degree of intelligence operating in a terrestrial environment?
- Any intelligent being?

Ideally, the researcher would begin the study with a clear idea of what the population of interest is, since the process of designing a study and testing hypotheses about the data that it produces does depend on the population about which you want to make statements. However, that doesn't always happen in practice: usually the researcher has a fairly vague idea of what the population is and designs the study as best he/she can on that basis. In practice this issue is often the source of very real headaches and very serious methodological worries.

10.1.1 Most samples are not simple random samples

To see where the concern comes from, it's important to recognise that – regardless of what the population actually is – whenever we're engaged in statistical inference, what we're trying to do is *generalise* from the sample to the population. In order to do so, we generally want to assume that the sample is drawn from the population “at random”. That is, in an ideal world we'd be able to say that every single member of the population is equally likely to end up being included in the sample. This is often referred to as a **simple random sample**. As you can see from looking at the list of possible populations that I showed above, it is almost impossible to obtain a simple random sample from most of the populations of interest. When I run experiments, I'd consider it a minor miracle if my participants turned out to be a random sampling of the undergraduate psychology students at Adelaide university, even though this is by far the narrowest population that I might want to generalise to.

Clearly, this can pose real difficulties in terms of external validity, as we discussed in Chapter 2. However, it's not quite as bad as it sounds. In truth, we don't need the sample to be randomly generated in *every* respect: we only need it to be random with respect to the psychologically-relevant phenomenon of interest. Suppose, for instance, I'm doing a study looking at working memory capacity. In study 1, I actually have the ability to sample randomly from all human beings currently alive, with one exception: I can only sample people born on a Monday. In study 2, I am able to sample randomly from the Australian population. I want to generalise my results to the population of all living humans. Which study is better? The answer, obviously, is study 1. Why? Because we have no reason to think that being “born on a Monday” has any interesting relationship to working memory capacity. In contrast, I can think of several reasons why “being Australian” might matter. Australia is a wealthy, industrialised country with a very well-developed education system. People growing up in that system will have had life experiences much more similar to the experiences of the people who designed the tests for working memory capacity. This shared experience might easily translate into similar beliefs about how to “take a test”, a shared assumption about how psychological experimentation works, and so on. These things might actually matter. For instance, “test taking” style might have taught the Australian participants how to direct their attention exclusively on fairly abstract test materials relative to people that haven't grown up in a similar environment; leading to a misleading picture of what working memory capacity is.

There are two points hidden in this discussion. Firstly, when designing your own studies, it's important to think about what population you care about, and try hard to sample in a way that is appropriate to that population. In practice, you're usually forced to put up with a “sample of convenience” (e.g., psychology lecturers sample psychology students because that's the least expensive way to collect data,

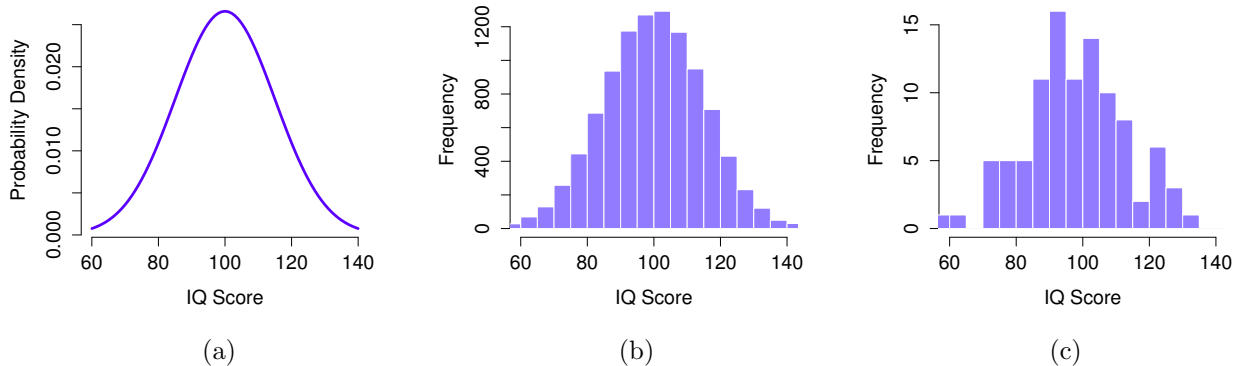


Figure 10.1: The population distribution of IQ scores (panel a) and two samples drawn randomly from it. In panel b we have a sample of 10,000 observations, and in panel c we have a sample of 100 observations

and our coffers aren't exactly overflowing with gold), but if so you should at least spend some time thinking about what the dangers of this practice might be.

Secondly, if you're going to criticise someone else's study because they've used a sample of convenience rather than laboriously sampling randomly from the entire human population, at least have the courtesy to offer a specific theory as to *how* this might have distorted the results. Remember, everyone in science is aware of this issue, and does what they can to alleviate it. Merely pointing out that "the study only included people from group BLAH" is entirely unhelpful, and borders on being insulting to the researchers, who are of course aware of the issue, they just don't happen to be in possession of the infinite supply of time and money required to construct the perfectly representative sample. In short, if you want to offer a responsible critique of the sampling process, then be *helpful*. Rehashing the blindingly obvious truisms that I've been rambling on about in this section isn't helpful.

10.1.2 Populations, samples and probability distributions

Okay. Setting aside the thorny methodological issues associated with obtaining a random sample and my rather unfortunate tendency to rant about lazy methodological criticism, let's consider a slightly different issue. In much the same way that psychological researchers need to operationalise our abstract theoretical ideas in terms of concrete measurements, statisticians need to operationalise the concept of a "population" in terms of mathematical objects that they know how to work with. You've already come across these objects: they're called probability distributions.

The idea is quite simple, really. Let's say we're talking about IQ scores, for example. By convention, the tests are designed so that, across the entire population, IQ scores follow a normal distribution that has mean 100 and standard deviation 15. As a statistical abstraction then, the "population" we're interested in is operationalised in terms of the probability distribution depicted in Figure 10.1a. To put this in formal mathematical language, this population distribution is normal, and described by two **population parameters**, the population mean $\mu = 100$ and the population standard deviation $\sigma = 15$.

Now let's suppose we conducted a massive experiment, with a **sample size** of $N = 10,000$ people, and measured their IQ. Let's also assume that the sampling is perfectly random. We can mimic this by using the `rnorm()` command, as discussed in the last chapter:

```
> X <- rnorm( n = 10000, mean = 100, sd = 15 )
```

Not surprisingly, since we have such a huge mass of data, when I draw a histogram of this sample, it looks very similar to the population distribution, as illustrated in Figure 10.1b. This data set can be characterised in terms of two **sample statistics**, the sample mean $\bar{X} = 99.89$ and the sample standard deviation $s = 15.07$. These are awfully close to the true values, although they're not quite perfect.

Now let's suppose that we collected a smaller sample, with only $N = 100$ people this time. If we draw the histogram for this sample (Figure 10.1c) it's pretty clear that that the sample does resemble the population, but it's not anywhere near as exact as the big sample. When we calculate the sample mean, we get a value of $\bar{X} = 98.45$; for the sample standard deviation, we obtain $s = 15.86$. We're now off by a bigger margin in both cases, though people are often surprised at just how close we get with a sample of only 100.

10.2

Estimating population means and standard deviations

In the IQ example in the previous section, we actually knew the population parameters ahead of time. As every undergraduate gets taught in their very first lecture on the measurement of intelligence, IQ scores are *defined* to have mean 100 and standard deviation 15. However, this is a bit of a lie. How do we know that IQ scores have a true population mean of 100? Well, we know this because the people who designed the tests have administered them to very large samples, and have then “rigged” the scoring rules so that their sample has mean 100. That’s not a bad thing of course: it’s an important part of designing a psychological measurement. However, it’s important to keep in mind that this theoretical mean of 100 only attaches to the population that the test designers used to design the tests. Good test designers will actually go to some lengths to provide “test norms” that can apply to lots of different populations (e.g., different age groups, nationalities etc).

This is very handy, but of course almost every research project of interest involves looking at a different population of people to those used in the test norms. For instance, suppose you wanted to measure the effect of low level lead poisoning on cognitive functioning in Port Pirie, a South Australian industrial town with a lead smelter. Perhaps you decide that you want to compare IQ scores among people in Port Pirie to a comparable sample in Whyalla, a South Australian industrial town with a steel refinery.¹ Regardless of which town you’re thinking about, it doesn’t make a lot of sense simply to *assume* that the true population mean IQ is 100. No-one has, to my knowledge, produced sensible norming data that can automatically be applied to South Australian industrial towns. We’re going to have to **estimate** the population means from a sample of data. So how do we do this?

For the sake of argument, suppose we go to Port Pirie, and 100 of the locals – sampled entirely at random, somehow – are kind enough to sit through an IQ test. Across these 100 people, the mean IQ score turned out to be $\bar{X} = 98.52$, let’s say. So, what is the true mean IQ μ for the population of Port Pirie?

¹Please note that if you were *actually* interested in this question, you would need to be a *lot* more careful than I’m being here. You *can’t* just compare IQ scores in Whyalla to Port Pirie and assume that any differences are due to lead poisoning. Even if it were true that the only differences between the two towns corresponded to the different refineries (and it isn’t, not by a long shot), you need to account for the fact that people already *believe* that lead pollution causes cognitive deficits: if you recall back to Chapter 2, this means that there are different demand effects for the Port Pirie sample than for the Whyalla sample. In other words, you might end up with an illusory group difference in your data, caused by the fact that people *think* that there is a real difference. I find it pretty implausible to think that the locals wouldn’t be well aware of what you were trying to do if a bunch of researchers turned up in Port Pirie with lab coats and IQ tests, and even less plausible to think that a lot of people would be pretty resentful of you for doing it. Those people won’t be as co-operative in the tests. Other people in Port Pirie might be *more* motivated to do well because they don’t want their home town to look bad. The motivational effects that would apply in Whyalla are likely to be weaker, because people don’t have any concept of “iron ore poisoning” in the same way that they have a concept for “lead poisoning”. Psychology is *hard*.

The correct answer here is that we don't actually know. Maybe it's 97.23, maybe it's 100.56. We can't say for sure. However, if I was forced at gunpoint to give a "best guess", I'd have to say 98.52. I don't have to believe that my guess is true (and I don't): all I believe here is that this is the best single number that I can give. So that's my estimate right there. It's not terribly complicated, since my estimate of the population mean is identical to the observed sample mean, but it's conceptually important. Statisticians often use the notation $\hat{\mu}$ to refer to "an estimate of μ ". So, if I observe a *sample mean* of $\bar{X} = 98.52$, then my **estimate of the population mean** is $\hat{\mu} = 98.52$. The distinction is subtle, but it's important. Just so that we're clear on the notation:

Symbol	What is it?	Do we know what it is?
\bar{X}	Sample mean	Yes, calculated from the raw data
μ	True population mean	Almost never known for sure
$\hat{\mu}$	Estimate of the population mean	Yes, identical to the sample mean

So far, estimation seems pretty simple. In the case of the mean, our estimate of the population parameter (i.e. $\hat{\mu}$) turned out to be identical to the corresponding sample statistic (i.e. \bar{X}). However, that's not always true. To see this, let's have a think about how to construct an **estimate of the population standard deviation**, which we'll denote $\hat{\sigma}$. What shall we use as our estimate in this case? Your first thought might be that we could do the same thing we did when estimating the mean, and just use the sample statistic as our estimate. That's almost the right thing to do, but not quite.

Here's why. Suppose I have a sample that contains a single observation. For this example, it helps to consider a sample where you have no intuitions at all about what the true population values might be, so let's use something completely fictitious. Suppose the observation in question measures the *cromulence* of my shoes. It turns out that my shoes have a cromulence of 20. So here's my sample:

20

This is a perfectly legitimate sample, even if it does have a sample size of $N = 1$. It has a sample mean of 20, and – because every observation in this sample is equal to the sample mean (obviously!) – it has a sample standard deviation of 0. Now, as a description of the *sample*, this seems quite right: the sample contains a single observation and therefore there is no variation observed within the sample. A sample standard deviation of $s = 0$ is the right answer here. But as an estimate of the *population* standard deviation, it feels completely insane, right? Admittedly, you and I don't know anything at all about what "cromulence" is, but we know something about data: the only reason that we don't see any variability in the *sample* is that the sample is too small to display any variation! So, if you have a sample size of $N = 1$, it *feels* like the right answer is just to say "no idea at all".

Notice that you *don't* have the same intuition when it comes to the sample mean and the population mean. If forced to make a best guess about the population mean, it doesn't feel completely insane to guess that the population mean is 20. Sure, you probably wouldn't feel very confident in that guess, because you have only the one observation to work with, but it's still the best guess you can make.

Let's extend this example a little. Suppose I now make a second observation. My data set now has $N = 2$ observations of the cromulence of shoes, and the complete sample now looks like this:

20, 22

This time around, our sample is *just* large enough for us to be able to observe some variability: two observations is the bare minimum number needed for any variability to be observed! For our new data set, the sample mean is $\bar{X} = 21$, and the sample standard deviation is $s = 1$. What intuitions do we have about the population? Again, as far as the population mean goes, the best guess we can possibly make is the sample mean: if forced to guess, we'd probably guess that the population mean cromulence is 21. What about the standard deviation? This is a little more complicated. The sample standard deviation

is only based on two observations, and if you're at all like me you probably have the intuition that, with only two observations, we haven't given the population "enough of a chance" to reveal its true variability to us. It's not just that we suspect that the estimate is *wrong*: after all, with only two observations we expect it to be wrong to some degree. The worry is that the error is *systematic*. Specifically, we suspect that the sample standard deviation is likely to be smaller than the population standard deviation.

This intuition feels right, but it would be nice to demonstrate this somehow. One way to do this would be to mathematically prove that the sample variance is systematically **biased** as an estimator of the population variance. That's what statisticians did originally, but unless you have the mathematical background it's not all that helpful. Instead, what I'll do is show you the results of some simulations done using R. Let's return to our IQ example. What I did was use the `rnorm()` function to generate 10,000 random data sets, each one consisting of a single observation (i.e., $N = 1$) drawn from a population that has a true mean of 100 and a true standard deviation of 15. I then did the same thing for data sets with a sample size of $N = 2$, and so on up to a sample size of $N = 10$. For every simulated data set, I calculated the sample mean and the sample standard deviation. Despite sounding like a pretty complicated number crunching exercise, it only takes a few seconds to run on a laptop. The results of this little exercise are shown in Figure 10.2. On the left hand side (panel a), I've plotted the average of the 10,000 simulated sample means, for all values N from 1 to 10. On the right hand side (panel b), I've plotted the average of the 10,000 standard deviations. The two plots are quite different: *on average*, the sample mean is equal to the population mean. It is an **unbiased estimator**, which is essentially the reason why your best estimate for the population mean is the sample mean.² The plot on the right is quite different: *on average*, the sample standard deviation s is *smaller* than the population standard deviation σ . It is a **biased estimator**. In other words, if we want to make a "best guess" $\hat{\sigma}$ about the value of the population standard deviation σ , we should make sure our guess is a little bit larger than the sample standard deviation s .

The fix to this systematic bias turns out to be very simple. Here's how it works. Before tackling the standard deviation, let's look at the variance. If you recall from Section 5.2, the sample variance is defined to be the average of the squared deviations from the sample mean. That is:

$$s^2 = \frac{1}{N} \sum_{i=1}^N (X_i - \bar{X})^2$$

The sample variance s^2 is a biased estimator of the population variance σ^2 . But as it turns out, we only need to make a tiny tweak to transform this into an unbiased estimator. All we have to do is divide by $N - 1$ rather than by N . If we do that, we obtain the following formula:

$$\hat{\sigma}^2 = \frac{1}{N - 1} \sum_{i=1}^N (X_i - \bar{X})^2$$

This is an unbiased estimator of the population variance σ . Moreover, this finally answers the question we raised in Section 5.2. Why did R give us slightly different answers when we used the `var()` function? Because the `var()` function calculates $\hat{\sigma}^2$ not s^2 , that's why. A similar story applies for the standard deviation. If we divide by $N - 1$ rather than N , our estimate of the population standard deviation becomes:

$$\hat{\sigma} = \sqrt{\frac{1}{N - 1} \sum_{i=1}^N (X_i - \bar{X})^2}$$

²I should note that I'm hiding something here. Unbiasedness is a desirable characteristic for an estimator, but there are other things that matter besides bias. However, it's beyond the scope of this book to discuss this in any detail. I just want to draw your attention to the fact that there's some hidden complexity here.

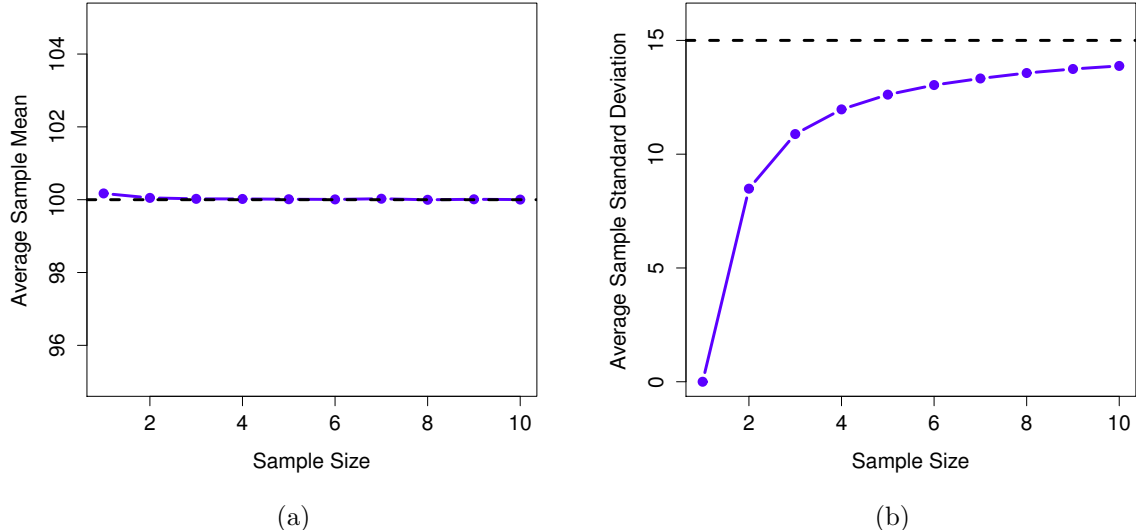


Figure 10.2: An illustration of the fact that the sample mean is an unbiased estimator of the population mean (panel a), but the sample standard deviation is a biased estimator of the population standard deviation (panel b). To generate the figure, I generated 10,000 simulated data sets with 1 observation each, 10,000 more with 2 observations, and so on up to a sample size of 10. Each data set consisted of fake IQ data: that is, the data were normally distributed with a true population mean of 100 and standard deviation 15. *On average*, the sample means turn out to be 100, regardless of sample size (panel a). However, the sample standard deviations turn out to be systematically too small (panel b), especially for small sample sizes.

and when we use R's built in standard deviation function `sd()`, what it's doing is calculating $\hat{\sigma}$, not s .³

One final point: in practice, a lot of people tend to refer to $\hat{\sigma}$ (i.e., the formula where we divide by $N - 1$) as the *sample* standard deviation. Technically, this is incorrect: the *sample* standard deviation should be equal to s (i.e., the formula where we divide by N). These aren't the same thing, either conceptually or numerically. One is a property of the sample, the other is an estimated characteristic of the population. However, in almost every real life application, what we actually care about is the estimate of the population parameter, and so people always report $\hat{\sigma}$ rather than s . This is the right number to report, of course, it's that people tend to get a little bit imprecise about terminology when they write it up, because "sample standard deviation" is shorter than "estimated population standard deviation". It's no big deal, and in practice I do the same thing everyone else does. Nevertheless, I think it's important to keep the two *concepts* separate: it's never a good idea to confuse "known properties of your sample" with "guesses about the population from which it came". The moment you start thinking that s and $\hat{\sigma}$ are the same thing, you start doing exactly that.

³Okay, I'm hiding something else here. In a bizarre and counterintuitive twist, since $\hat{\sigma}^2$ is an unbiased estimator of σ^2 , you'd assume that taking the square root would be fine, and $\hat{\sigma}$ would be an unbiased estimator of σ . Right? Weirdly, it's not. There's actually a subtle, tiny bias in $\hat{\sigma}$. This is just bizarre: $\hat{\sigma}^2$ is an unbiased estimate of the population variance σ^2 , but when you take the square root, it turns out that $\hat{\sigma}$ is a biased estimator of the population standard deviation σ . Weird, weird, weird, right? So, why is $\hat{\sigma}$ biased? The technical answer is "because non-linear transformations (e.g., the square root) don't commute with expectation", but that just sounds like gibberish to everyone who hasn't taken a course in mathematical statistics. Fortunately, it doesn't matter for practical purposes. The bias is small, and in real life everyone uses $\hat{\sigma}$ and it works just fine. Sometimes mathematics is just annoying.

To finish this section off, here's another couple of tables to help keep things clear:

Symbol	What is it?	Do we know what it is?
s	Sample standard deviation	Yes, calculated from the raw data
σ	Population standard deviation	Almost never known for sure
$\hat{\sigma}$	Estimate of the population standard deviation	Yes, but not the same as the sample standard deviation
Symbol	What is it?	Do we know what it is?
s^2	Sample variance	Yes, calculated from the raw data
σ^2	Population variance	Almost never known for sure
$\hat{\sigma}^2$	Estimate of the population variance	Yes, but not the same as the sample variance

10.3

Sampling distributions

When talking about probability distributions, we've made use of the idea that observations X are sampled from some distribution. So, for instance, if X refers to a measurement of an IQ score, then we would say that the distribution of X is normal, with mean $\mu = 100$ and standard deviation $\sigma = 15$. If I repeated it 10,000 times, and drew a histogram of all the individual X values that I measured, I'd end up with a histogram that looks like the one in Figure 10.3a.

This idea can be extended, to talk about the distribution of a sample statistic. That is, suppose I gave $N = 10$ people an IQ test, and calculated the mean IQ for that sample \bar{X} . It's perfectly reasonable to think about the sample mean itself \bar{X} as being a kind of measurement. If I repeated this procedure 100,000 times, and drew a histogram of all the sample means \bar{X} that I measured, what would I end up with? If I use R to simulate what would happen, I end up with Figure 10.3c. Obviously, since we can draw a histogram of these \bar{X} values, we should be able to talk about \bar{X} as having a distribution, right? After all, by plotting the histogram, we can clearly see what the distribution is! The name for this distribution over \bar{X} is the **sampling distribution of the mean**, and you can see from this plot that there's a relationship between the population distribution (i.e., the distribution of X) and the sampling distribution of the mean (i.e., the distribution of \bar{X}).

10.4

The central limit theorem

In the example above, the sampling distribution looks like it's a normal distribution with the same mean as our original population distribution (100), but a narrower standard deviation. This narrowing is important, and it makes a lot of sense: we're looking at the distribution of the sample mean \bar{X} , and you'd kind of expect a the sample mean to be closer the population mean, right? That's why we collect lots of data in the first place! What you're looking at in this picture is exactly this intuition in action: the sample means \bar{X} all tend to "cluster" around the true mean μ , much more tightly than a single observation X would be expected to do. This phenomenon is an illustration of the **law of large numbers**. As the sample size increases, it becomes more and more likely that your sample mean will be very close to the true answer. Or to be precise, what the law of large numbers says is that, as the sample size approaches infinity (written as $N \rightarrow \infty$) the sample mean approaches the population mean ($\bar{X} \rightarrow \mu$).

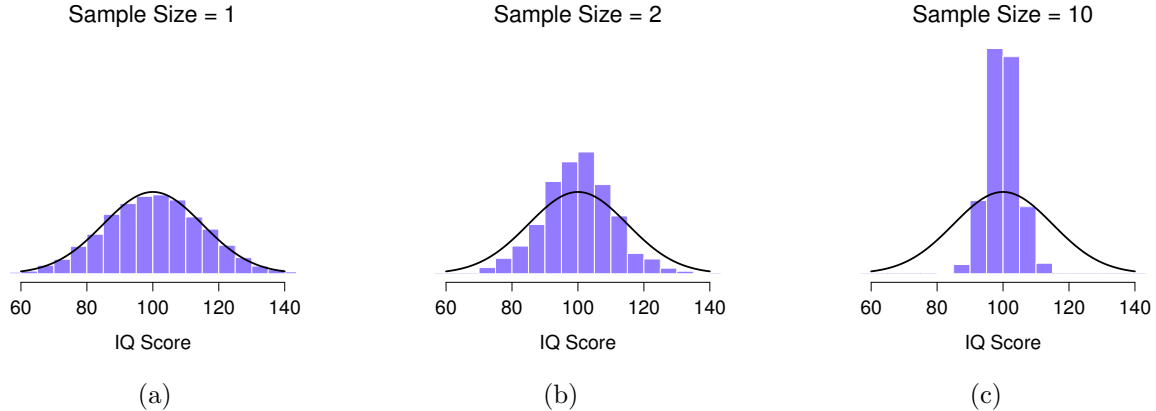


Figure 10.3: An illustration of the sampling distribution of the mean. In each panel, I generated 10,000 samples of IQ data, and calculated the mean IQ observed within each of these data sets. The histograms in these plots show the distribution of these means (i.e., the sampling distribution of the mean). Each individual IQ score was drawn from a normal distribution with mean 100 and standard deviation 15, which is shown as the solid black line). In panel a, each data set contained only a single observation, so the mean of each sample is just one person's IQ score. As a consequence, the sampling distribution of the mean is of course identical to the population distribution of IQ scores. However, when we raise the sample size to 2, the mean of any one sample tends to be closer to the population mean than a one person's IQ score, and so the histogram (i.e., the sampling distribution) is a bit narrower than the population distribution. By the time we raise the sample size to 10 (panel c), we can see that the distribution of sample means tend to be fairly tightly clustered around the true population mean.

.....

The law of large numbers is a very powerful tool, but we can actually make a stronger statement than this, using one of the most useful mathematical results in all of statistics: the **central limit theorem**. Let's suppose that the population distribution of X has a mean μ and standard deviation σ , but it's *not* a normal distribution. The central limit theorem says that, as the sample size increases, the sampling distribution of the mean \bar{X} will *become* a normal distribution, with mean μ and standard deviation σ/\sqrt{N} . By convention, the standard deviation of a sampling distribution is called the **standard error** (SE). Therefore, the standard error of the mean \bar{X} , can be described by the following equation,

$$\text{SE}(\bar{X}) = \frac{\sigma}{\sqrt{N}}$$

Hm. While I'll concede that the standard error of the mean is actually a very useful thing to be able to calculate, I feel that this equation is overselling itself slightly.⁴

Be that as it may, we should probably get back to the stats... the central limit theorem is a pretty important thing in statistics, so I'll "prove" it to you in a moment, but before I do so, let's just take a moment to think about just how incredible a claim this is. There's a *lot* of variables that you might want to measure that could be thought of as being the average of lots of different things. What the central limit theorem implies is that, whenever that is the case, you should expect your variable to follow a normal distribution. And remarkably, this really happens. Over and over again in the real world you find that variables you care about are approximately normally distributed. At a fundamental level, the

⁴Sure, I *could* have chosen "SEM" as the notation for the standard error of the mean. I didn't do this for two reasons: (1) later on we'll want to talk about the standard errors for things other than the mean, so it's better to be able to use notation that will handle this nicely, and (2) it's funnier this way.

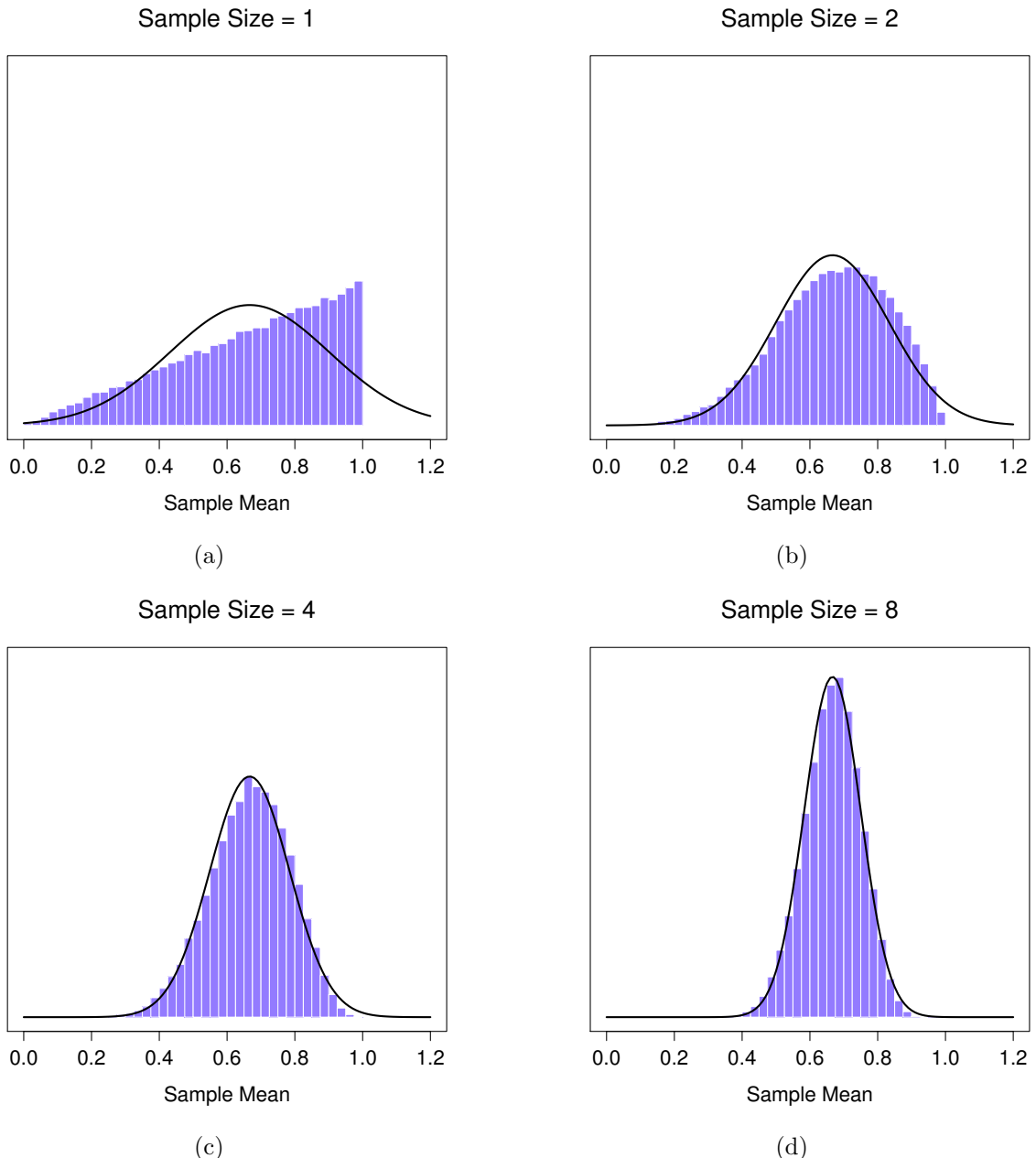


Figure 10.4: A demonstration of the central limit theorem. In panel a, we have a non-normal population distribution; and panels b-d show the sampling distribution of the mean for samples of size 2,4 and 8, for data drawn from the distribution in panel a. As you can see, even though the original population distribution is non-normal, the sampling distribution of the mean becomes pretty close to normal by the time you have a sample of even 4 observations.

central limit theorem is the mathematical reason why.

Okay, time to demonstrate the central limit theorem in action. I'm not going to give a mathematical proof, since very few people would want to read it. Instead, I'm going to do the same thing I've done throughout this chapter: I'll use R to simulate it. Let's start out with a very non-normal population, namely the "ramped" distribution shown in Figure 10.4a. This plot shows a histogram of 100,000 samples, where I've plotted the appropriate normal distribution over the top to give you a sense of what to compare it to. Next, let's see what happens then if we generate 100,000 samples of size $N = 2$, and then plot the histogram of the means of these samples? This time, we get the triangular distribution shown in Figure 10.4b. That's not normal either, but it's definitely closer. When, we increase the sample size to $N = 3$, the sampling distribution of the mean looks closer to normal, as illustrated in Figure 10.4c, and by the time we've raised our sample size to a massive $N = 4$, as shown in Figure 10.4, it's pretty hard to think that this is anything other than a normal distribution. In other words, while the central limit theorem *technically* refers only to things getting normal as N approaches infinity, in this example all it took was a sample size of 4. Which is a bit smaller than infinity, I must admit. The take home message is that everything turns into a normal distribution, eventually.⁵

10.5

Estimating a confidence interval

The last thing I'm going to talk about in this chapter are confidence intervals. The estimates that we talked about in the last section are all examples of "point" estimates. What I mean by that is that what we're doing is making a single "best guess" about what the value of a particular population parameter is. Most of the time, we don't just want a single best guess, we also want to be able to estimate a range of values, in such a way that we can feel pretty confident that the range includes the true value. The name for this is a **confidence interval**.

This is actually pretty easy, since all the hard work was done in the previous sections. To see how this works, let's suppose for the moment that we actually knew the true mean μ and the true standard deviation σ . Then we would also be able to say that (as long as N is big enough) the sampling distribution for the mean \bar{X} of a sample drawn from this population would be normal with mean μ . Not only that, we also know that the standard error of the mean is calculated by dividing the population standard deviation σ by the square root of the sample size \sqrt{N} . Now, remember from our previous discussion of the normal distribution in Section 9.4 that 95% of the distribution lies within 2 standard deviations of the mean? To be slightly more precise, we can use the `qnorm()` function to compute the 2.5th and 97.5th percentiles of the normal distribution

```
> qnorm( p = c(.025, .975) )
[1] -1.959964  1.959964
```

and we see that in fact 95% of the distribution falls within 1.96 standard deviations either side of the mean. What does that tell us about \bar{X} ? Well, it tells us that 95% of all data sets (of size N) that we could sample from this population will have a sample mean \bar{X} that falls within 1.96 standard errors of the mean! In other words, there is a 95% probability that:

$$\mu - \left(1.96 \times \frac{\sigma}{\sqrt{N}} \right) \leq \bar{X} \leq \mu + \left(1.96 \times \frac{\sigma}{\sqrt{N}} \right)$$

⁵Well, sort of. The central limit theorem doesn't cover every possible situation, but it is very very broad. Like most introductory stats texts, I've discussed one situation where the central limit theorem holds: when you're taking an average across lots of independent events drawn from the same distribution. However, the central limit theorem is much broader than this. There's a whole class of things called "U-statistics" for instance, all of which satisfy the central limit theorem and therefore become normally distributed for large sample sizes. The mean is one such statistic, but it's not the only one.

Okay, that's all well and good, but in some ways it's the opposite of what we're interested in. The equation above tells us what we should expect about the sample mean, given that we know what the population parameters are. What we want is to have this work the other way around: we want to know what we should believe about the population parameters, given that we have observed a particular sample. However, it's not too difficult to do this. Using a little high school algebra, a sneaky way to rewrite our equation is like this:

$$\bar{X} - \left(1.96 \times \frac{\sigma}{\sqrt{N}} \right) \leq \mu \leq \bar{X} + \left(1.96 \times \frac{\sigma}{\sqrt{N}} \right)$$

What this is telling us is that the range of values has a 95% probability of containing the population mean μ . We refer to this range as a **95% confidence interval**, denoted CI_{95} . In short, as long as N is sufficiently large – large enough for us to believe that the sampling distribution of the mean is normal – then we can write this as our formula for the confidence interval:

$$CI_{95} = \bar{X} \pm \left(1.96 \times \frac{\sigma}{\sqrt{N}} \right)$$

Of course, there's nothing terribly special about the value 1.96, other than the fact it's the number of standard deviations away from the mean that you need to extend your interval to cover 95% of the sampling distribution. If I'd wanted a 70% confidence interval, I could have used the `qnorm()` function to calculate the 15th and 85th quantiles:

```
> qnorm( p = c(.15, .85) )
[1] -1.036433  1.036433
```

and so the formula for CI_{70} would be the same as the formula for CI_{95} except that we'd use 1.04 as our magic number rather than 1.96.

Unfortunately, this formula assumes that we actually know the true population standard deviation σ . In practice, we never really do know this, so we have to use an estimate of the standard deviation $\hat{\sigma}$ instead. This is pretty straightforward to do, but (for reasons we'll talk about in Chapter 13) this has the consequence that we need to use the quantiles of the t -distribution rather than the normal distribution to calculate our magic number; and the answer depends on the sample size. When N is very large, we get pretty much the same value using `qt()` that we would if we used `qnorm()`...

```
> N <- 10000  # suppose our sample size is 10,000
> qt( p = .975, df = N-1)  # calculate the 97.5th quantile of the t-dist
[1] 1.960201
```

But when N is small, we get a much bigger number when we use the t distribution:

```
> N <- 10  # suppose our sample size is 10
> qt( p = .975, df = N-1)  # calculate the 97.5th quantile of the t-dist
[1] 2.262157
```

There's nothing too mysterious about what's happening here. Bigger values mean that the confidence interval is wider, indicating that we're more uncertain about what the true value of μ actually is. When we use the t distribution instead of the normal distribution, we get bigger numbers, indicating that we have more uncertainty. And why do we have that extra uncertainty? Well, because our estimate of the population standard deviation $\hat{\sigma}$ might be wrong! If it's wrong, it implies that we're a bit less sure about what our sampling distribution of the mean actually looks like... and this uncertainty ends up getting reflected in a wider confidence interval.

10.5.1 Calculating confidence intervals in R

As far as I can tell, the core packages in R don't include a simple function for calculating confidence intervals for the mean. They *do* include a lot of complicated, extremely powerful functions that can be used to calculate confidence intervals associated with lots of different things, such as the `confint()` function that we'll use in Chapter 15. But I figure that when you're first learning statistics, it might be useful to start with something simpler. As a consequence, the `lsr` package includes a function called `ciMean()` which you can use to calculate your confidence intervals. There are two arguments that you might want to specify:⁶

- `x`. This should be a numeric vector containing the data.
- `conf`. This should be a number, specifying the confidence level. By default, `conf = .95`, since 95% confidence intervals are the de facto standard in psychology.

So, for example, if I load the `afl24.Rdata` file, calculate the confidence interval associated with the mean attendance:

```
> ciMean( x = afl$attendance )
  2.5%    97.5%
31597.32 32593.12
```

Hopefully that's fairly clear.

10.5.2 Plotting confidence intervals in R

There's several different ways you can draw graphs that show confidence intervals as error bars. I'll show three versions here, but this certainly doesn't exhaust the possibilities. In doing so, what I'm assuming is that you want to draw is a plot showing the means and confidence intervals for one variable, broken down by different levels of a second variable. For instance, in our `afl` data that we discussed earlier, we might be interested in plotting the average `attendance` by `year`. I'll do this using three different functions, `bargraph.CI()`, `lineplot.CI()` (both of which are in the `sciplot` package), and `plotmeans()` (which is in the `gplots` package). Assuming that you've installed these packages on your system (see Section 4.2 if you've forgotten how to do this), you'll need to load them. You'll also need to load the `lsr` package, because we'll make use of the `ciMean()` function to actually calculate the confidence intervals

```
> library( sciplot )      # bargraph.CI() and lineplot.CI() functions
> library( gplots )       # plotmeans() function
> library( lsr )          # ciMean() function
```

Here's how to plot the means and confidence intervals drawn using `bargraph.CI()`.

```
> bargraph.CI( x.factor = year,           # grouping variable
+               response = attendance,     # outcome variable
+               data = afl,                # data frame with the variables
+               ci.fun= ciMean,            # name of the function to calculate CIs
+               xlab = "Year",              # x-axis label
+               ylab = "Average Attendance" # y-axis label
+ )
```

⁶As of the current writing, these are the only arguments to the function. However, I am planning to add a bit more functionality to `ciMean()`. However, regardless of what those future changes might look like, the `x` and `conf` arguments will remain the same, and the commands used in this book will still work.

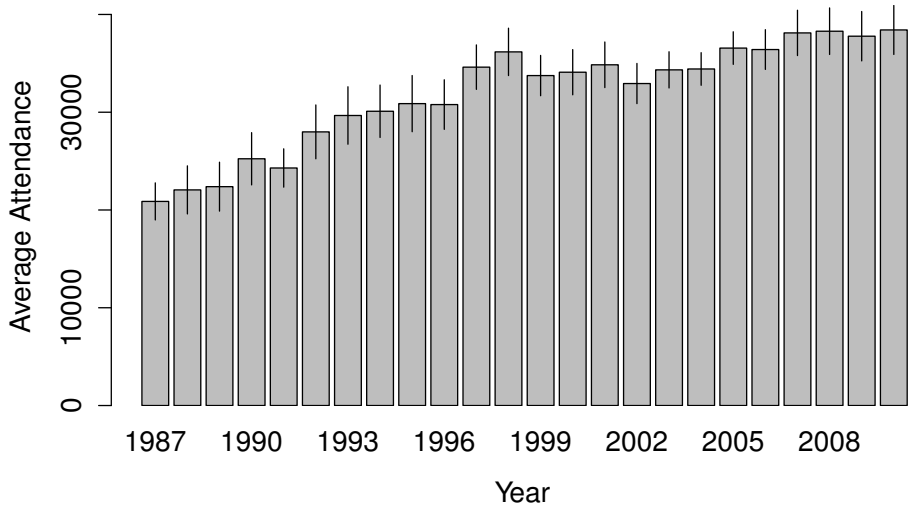


Figure 10.5: Means and 95% confidence intervals for AFL `attendance`, plotted separately for each `year` from 1987 to 2010. This graph was drawn using the `bargraph.CI()` function.

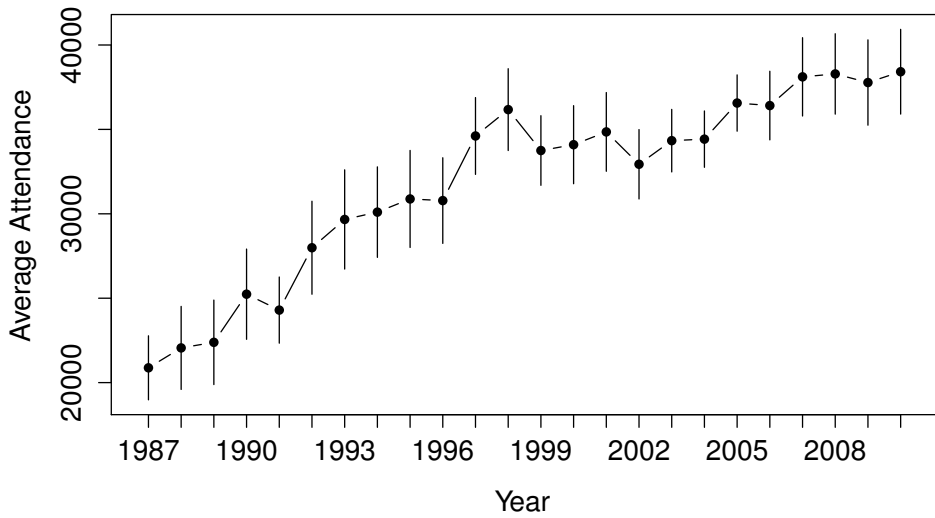


Figure 10.6: Means and 95% confidence intervals for AFL `attendance`, plotted separately for each `year` from 1987 to 2010. This graph was drawn using the `lineplot.CI()` function.

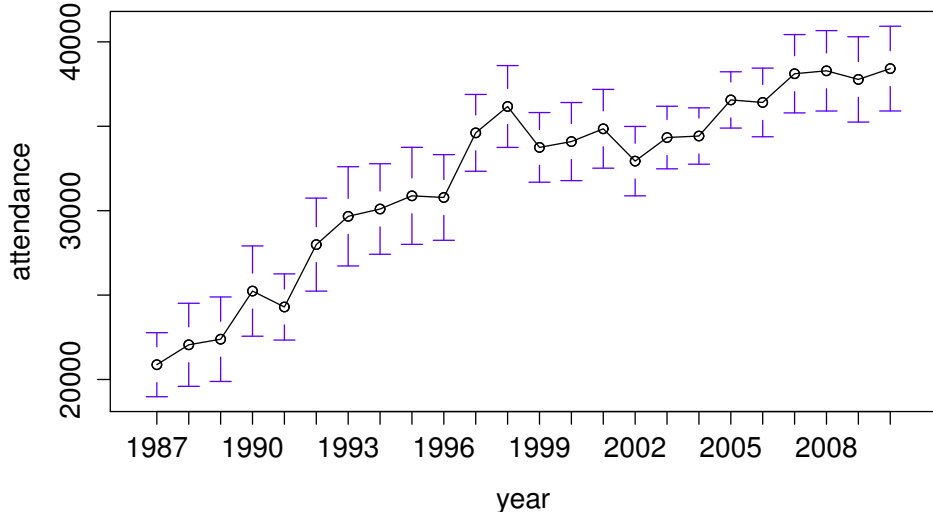


Figure 10.7: Means and 95% confidence intervals for AFL `attendance`, plotted separately for each `year` from 1987 to 2010. This graph was drawn using the `plotmeans()` function.

This produces the output shown in Figure 10.5. We can use the same arguments when calling the `lineplot.CI()` function:

```
> lineplot.CI( x.factor = year,           # grouping variable
+               response = attendance,   # outcome variable
+               data = afl,              # data frame with the variables
+               ci.fun= ciMean,          # name of the function to calculate CIs
+               xlab = "Year",            # x-axis label
+               ylab = "Average Attendance" # y-axis label
+ )
```

And the output for this command is shown in Figure 10.6. Finally, here's how you would do it using `plotmeans()`:

```
> plotmeans( formula = attendance ~ year, # outcome ~ group
+             data = afl,                  # data frame with the variables
+             n.label = FALSE              # don't show the sample sizes
+ )
```

This is shown in Figure 10.7.

10.5.3 Interpreting a confidence interval

The most counterintuitive thing about confidence intervals is how they are supposed to be interpreted. Whenever people first encounter confidence intervals, their first instinct is almost always to interpret it

like this:

There is a 95% chance that the population mean falls within the 95% confidence interval.

This is the natural, intuitive and obvious way to think about the confidence interval. Strictly speaking, however, you're not allowed to do this. Remember how, back in Section 9.6, I said that there were two warring theories of what the word "probability" means? And I said that it would turn out to matter some time? Well, that time is now.

The basic problem with the statement that I made above is that you're making a probabilistic statement about the true population mean (i.e., you're saying that there's a 95% chance that the population mean takes on a value that fall within some fixed range). However, confidence intervals are a tool invented within the frequentist paradigm, and the frequentist definition of probability forbids this. Remember, the frequentist definition of probability requires that replication is possible. Because, in order for a probability statement to be meaningful, you have to be able to imagine "repeating the procedure" over and over again. This works really well for some things: for instance, I *can* say that "the probability of a coin flip being heads is 0.5", because I can imagine flipping a coin millions of times. And if I did flip a coin a million times, I'd expect about half of those coin flips to land on heads. Cool.

Okay, now let's try this on for size: "there is a 95% probability that the average global temperature is between 15 and 20 degrees" (or whatever). According to the frequentist view, this is gibberish. I can't generate a million planet Earths and measure their temperature. Therefore, there's no way to assign a probability value to "global average temperatures". In general, frequentists *aren't* allowed to make probability claims about population parameters. For frequentists, probabilities attach only to estimates, not to parameters.

The solution that we adopt is to rewrite the statement so that we're making a probability claim about the interval itself, *not* the population mean. If we say something like this

95% of all confidence intervals constructed using this procedure will include the corresponding population mean

we're being clear about the fact that we're only making probability statements about the confidence interval, and not about the population mean. Stating things this way is kind of awkward, and in practice no-one ever uses the explicitly frequentist statement. In fact, most people are quite happy to ignore the whole issue. But there are some people who get a bit pedantic about how confidence intervals should be interpreted, so as a rough guide it's helpful to think of it like this:

- "There is a **95% chance that a 95% confidence interval** will include the population mean." This statement seems to imply that the "95% chance" attaches to the confidence interval, so this is okay.
- "There is a **95% chance that the population mean** falls within the 95% CI." This statement seems to imply that the "95% chance" actually relates to the population mean. Occasionally people will ask you to rephrase this.

Kind of silly, isn't it?⁷

⁷As it happens, there's a magic wand that we can wave over the confidence interval to make this idiotic problem go away. It's called "Bayesian probability". All you have to do is be willing to say that probabilities are "in the mind" and not "in the world", and this whole problem vanishes. And in fact, Bayesians have a nearly identical tool called a *credible interval* which behaves almost the same way as a confidence interval, except that you're allowed to interpret it the natural way.

Summary

- Basic ideas about sampling (Section 10.1)
- Estimating means and standard deviations (Section 10.2)
- Sampling distributions for the mean and other statistics (Section 10.3)
- The central limit theorem (Section 10.4).
- Estimating a confidence interval (Section 10.5)

11. Hypothesis testing

NAPOLEON: *M. Laplace, they tell me you have written this large book on the system of the universe, and have never even mentioned its Creator.*

LAPLACE: *I had no need of that hypothesis.*

In the last chapter, I discussed the ideas behind estimation, which is one of the two “big ideas” in inferential statistics. It’s now time to turn our attention to the other big idea, which is *hypothesis testing*. In its most abstract form, hypothesis testing is a very simple idea: the researcher has some theory about the world, and wants to determine whether or not the data actually support that theory. However, the details are messy, and most people find the theory of hypothesis testing to be the most frustrating part of statistics. The structure of the chapter is as follows. Firstly, I’ll describe how hypothesis testing works, in a fair amount of detail, using a simple running example to show you how a hypothesis test is “built”. I’ll try to avoid being too dogmatic while doing so, and focus instead on the underlying logic of the testing procedure.¹ Afterwards, I’ll spend a bit of time talking about the various dogmas, rules and heresies that surround the theory of hypothesis testing.

11.1

A menagerie of hypotheses

In Umberto Eco’s book *Foucault’s Pendulum*, one of the main characters expounds a theory of the world in which all people can be classified as fools, morons, cretins and lunatics.² The lunatic category is the most awesome: I’m paraphrasing here because I’m too lazy to dig up the original...

You can tell him by his vaulting leaps of logic, that for him everything proves everything else,
and the fact that eventually it all comes down the Knights Templar

Personally I think we all succumb to lunacy eventually. For me, that day will arrive once I’m finally promoted to full professor (the last promotion I could possibly achieve, since everything above that counts as management; and I do *not* have managerial potential). Safely ensconced in my ivory tower, happily protected by tenure, I will finally be able to take leave of my senses (so to speak), and indulge in

¹A technical note. The description below differs subtly from the standard description given in a lot of introductory texts. The orthodox theory of null hypothesis testing emerged from the work of Sir Ronald Fisher and Jerzy Neyman in the early 20th century; but Fisher and Neyman actually had very different views about how it should work. The standard treatment of hypothesis testing that most texts use is a hybrid of the two approaches. The treatment here is a little more Neyman-style than the orthodox view, especially as regards the meaning of the *p* value.

²Feel free to indulge in speculation as to which one I am.

that most thoroughly unproductive line of psychological research: the search for extrasensory perception (ESP).³

Let's suppose that this glorious day has come. My first study is a simple one, in which I seek to test whether clairvoyance exists. Each participant sits down at a table, and is shown a card by an experimenter. The card is black on one side and white on the other. The experimenter takes the card away, and places it on a table in an adjacent room. The card is placed black side up or white side up completely at random, with the randomisation occurring only after the experimenter has left the room with the participant. A second experimenter comes in and asks the participant which side of the card is now facing upwards. It's purely a one-shot experiment. Each person sees only one card, and gives only one answer; and at no stage is the participant actually in contact with someone who knows the right answer. My data set, therefore, is very simple. I have asked the question of N people, and some number X of these people have given the correct response. To make things concrete, let's suppose that I have tested $N = 100$ people, and $X = 62$ of these got the answer right... a surprisingly large number, sure, but is it large enough for me to feel safe in claiming I've found evidence for ESP? This is the situation where hypothesis testing comes in useful. However, before we talk about how to *test* hypotheses, we need to be clear about what we mean by hypotheses.

11.1.1 Research hypotheses versus statistical hypotheses

The first distinction that you need to keep clear in your mind is between research hypotheses and statistical hypotheses. In my ESP study, my overall scientific goal is to demonstrate that clairvoyance exists. In this situation, I have a clear research goal: I am hoping to discover evidence for ESP. In other situations I might actually be a lot more neutral than that, so I might say that my research goal is to determine whether or not clairvoyance exists. Regardless of how I want to portray myself, the basic point that I'm trying to convey here is that a research hypothesis involves making a substantive, testable scientific claim... if you are a psychologist, then your research hypotheses are fundamentally *about* psychological constructs. Any of the following would count as **research hypotheses**:

- *Listening to music reduces your ability to pay attention to other things.* This is a claim about the causal relationship between two psychologically meaningful concepts (listening to music and paying attention to things), so it's a perfectly reasonable research hypothesis.
- *Intelligence is related to personality.* Like the last one, this is a relational claim about two psychological constructs (intelligence and personality), but the claim is weaker: correlational not causal.
- *Intelligence is speed of information processing.* This hypothesis has a quite different character: it's not actually a relational claim at all. It's an ontological claim about the fundamental character of intelligence (and I'm pretty sure it's wrong). It's worth expanding on this one actually: It's usually easier to think about how to construct experiments to test research hypotheses of the form "does X affect Y?" than it is to address claims like "what is X?" And in practice, what usually happens is that you find ways of testing relational claims that follow from your ontological ones. For instance, if I believe that intelligence *is* speed of information processing in the brain, my experiments will often involve looking for relationships between measures of intelligence and measures of speed. As a consequence, most everyday research questions do tend to be relational in nature, but they're almost always motivated by deeper ontological questions about the state of nature.

³My apologies to anyone who actually believes in this stuff, but on my reading of the literature on ESP, I feel relatively confident in concluding that ESP research is a litany of false positives (referred to as Type I errors). To be fair, though, some of the studies are rigorously designed; so it's actually an interesting area for thinking about psychological research design. And of course it's a free country, so you can spend your own time and effort proving me wrong if you like, but I wouldn't think that's a terribly practical use of your intellect.

Notice that in practice, my research hypotheses could overlap a lot. My ultimate goal in the ESP experiment might be to test an ontological claim like “ESP exists”, but I might operationally restrict myself to a narrower hypothesis like “Some people can ‘see’ objects in a clairvoyant fashion”. That said, there are some things that really don’t count as proper research hypotheses in any meaningful sense:

- *Love is a battlefield.* This is too vague to be testable. While it’s sort of okay for a research hypothesis to have a degree of vagueness to it, it has to be possible to operationalise your theoretical ideas. Maybe I’m just not creative enough to see it, but I can’t see how this can be converted into any concrete research design. And if that’s true, then this isn’t a scientific research hypothesis, it’s a pop song. That doesn’t mean it’s not interesting – a lot of deep questions that humans have fall into this category. Maybe one day science will be able to construct testable theories of love, or to test to see if God exists, and so on; but right now we can’t, and I wouldn’t bet on ever seeing a satisfying scientific approach to either.
- *The first rule of tautology club is the first rule of tautology club.* This is not a substantive claim of any kind. It’s true by definition. No conceivable state of nature could possibly be inconsistent with this claim. As such, we say that this is an unfalsifiable hypothesis, and as such it is outside the domain of science. Whatever else you do in science, your claims must have the possibility of being wrong.
- *More people in my experiment will say “yes” than “no”.* This one fails as a research hypothesis because it’s a claim about the data set, not about the psychology (unless of course your actual research question is whether people have some kind of “yes” bias!). As we’ll see shortly, this hypothesis is starting to sound more like a statistical hypothesis than a research hypothesis.

As you can see, research hypotheses can be somewhat messy at times; and ultimately they are *scientific* claims. **Statistical hypotheses** are neither of these two things. Statistical hypotheses must be mathematically precise, and they must correspond to specific claims about the characteristics of the data generating mechanism (i.e., the “population”). Even so, the intent is that statistical hypotheses bear a clear relationship to the substantive research hypotheses that you care about! For instance, in my ESP study my research hypothesis is that some people are able to see through walls or whatever. What I want to do is to “map” this onto a statement about how the data were generated. So let’s think about what that statement would be. The quantity that I’m interested in within the experiment is $P(\text{"correct"})$, the true-but-unknown probability with which the participants in my experiment answer the question correctly. Let’s use the Greek letter θ (theta) to refer to this probability. If ESP doesn’t exist, and if my experiment is well designed, then the true probability will be $\theta = 0.5$... if people are just guessing, they’ll get it right half of the time. Alternatively, suppose ESP does exist and people can see the card: if so, people will perform better than chance, so we would expect that $\theta > 0.5$. Or, as another alternative, suppose ESP does exist, but the colours are all reversed and people don’t realise it (okay, that’s wacky, but you never know...). If that’s how it works then you’d actually expect performance to be below chance. This would correspond to a statistical hypothesis that $\theta < 0.5$. Finally, suppose ESP exists, but I have no idea whether or not the colours are reversed. In that case, the only claim I could make about the data would be that the probability of making the correct answer is *not* equal to 50%, which we would write as $\theta \neq 0.5$. All of these are legitimate examples of a statistical hypothesis.

All of this is by way of saying that, when attempting to construct a statistical hypothesis test, the researcher actually has two quite distinct hypotheses to consider. First, he or she has a research hypothesis (a claim about psychology), and this corresponds to a statistical hypothesis (a claim about the data generating population). In my ESP example, these might be

$$\begin{aligned} \text{Dan's research hypothesis: } & \text{“ESP exists”} \\ \text{Dan's statistical hypothesis: } & \theta \neq 0.5 \end{aligned}$$

And the key thing to recognise is this: *a statistical hypothesis test is a test of the statistical hypothesis, not the research hypothesis*. If your study is badly designed, then the link between your research hypothesis and your statistical hypothesis is broken. To give a silly example, suppose that my ESP study was conducted in a situation where the participant can actually see the card reflected in a window; if that happens, I would be able to find very strong evidence that $\theta \neq 0.5$, but this would tell us nothing about whether “ESP exists”.

11.1.2 Null hypotheses and alternative hypotheses

So far, so good. I have a research hypothesis that corresponds to what I want to believe about the world, and I can map it onto a statistical hypothesis that corresponds to what I want to believe about how the data were generated. It’s at this point that things get somewhat counterintuitive for a lot of people. Because what I’m about to do is invent a new statistical hypothesis (the “null” hypothesis, H_0) that corresponds to the exact opposite of what I want to believe, and then focus exclusively on that, almost to the neglect of the thing I’m actually interested in (which is now called the “alternative” hypothesis, H_1). In our ESP example, the null hypothesis is that $\theta = 0.5$, since that’s what we’d expect if ESP didn’t exist. My hope, of course, is that ESP is totally real, and so the *alternative* to this null hypothesis is $\theta \neq 0.5$. In essence, what we’re doing here is dividing up the possible values of θ into two groups: those values that I really hope aren’t true (the null), and those values that I’d be happy with if they turn out to be right (the alternative). Having done so, the important thing to recognise is that the goal of a hypothesis test is *not* to show that the alternative hypothesis is (probably) true; the goal is to show that the null hypothesis is (probably) false. Most people find this pretty weird.

The best way to think about it, in my experience, is to imagine that a hypothesis test is a criminal trial⁴... *the trial of the null hypothesis*. The null hypothesis is the defendant, the researcher is the prosecutor, and the statistical test itself is the judge. Just like a criminal trial, there is a presumption of innocence: the null hypothesis is *deemed* to be true unless you, the researcher, can prove beyond a reasonable doubt that it is false. You are free to design your experiment however you like (within reason, obviously!), and your goal when doing so is to maximise the chance that the data will yield a conviction... for the crime of being false. The catch is that the statistical test sets the rules of the trial, and those rules are designed to protect the null hypothesis – specifically to ensure that if the null hypothesis is actually true, the chances of a false conviction are guaranteed to be low. This is pretty important: after all, the null hypothesis doesn’t get a lawyer. And given that the researcher is trying desperately to prove it to be false, *someone* has to protect it.

11.2

Two types of errors

Before going into details about how a statistical test is constructed, it’s useful to understand the philosophy behind it. I hinted at it when pointing out the similarity between a null hypothesis test and a criminal trial, but I should now be explicit. Ideally, we would like to construct our test so that we never make any errors. Unfortunately, since the world is messy, this is never possible. Sometimes you’re just really unlucky: for instance, suppose you flip a coin 10 times in a row and it comes up heads all 10 times. That feels like very strong evidence that the coin is biased (and it is!), but of course there’s a 1 in 1024 chance that this would happen even if the coin was totally fair. In other words, in real life we *always* have to accept that there’s a chance that we did the wrong thing. As a consequence, the goal behind statistical hypothesis testing is not to *eliminate* errors, but to *minimise* them.

⁴This analogy only works if you’re from an adversarial legal system like UK/US/Australia. As I understand these things, the French inquisitorial system is quite different.

At this point, we need to be a bit more precise about what we mean by “errors”. Firstly, let’s state the obvious: it is either the case that the null hypothesis is true, or it is false; and our test will either reject the null hypothesis or retain it.⁵ So, as the table below illustrates, after we run the test and make our choice, one of four things might have happened:

	retain H_0	reject H_0
H_0 is true	correct decision	error (type I)
H_0 is false	error (type II)	correct decision

As a consequence there are actually *two* different types of error here. If we reject a null hypothesis that is actually true, then we have made a **type I error**. On the other hand, if we retain the null hypothesis when it is in fact false, then we have made a **type II error**.

Remember how I said that statistical testing was kind of like a criminal trial? Well, I meant it. A criminal trial requires that you establish “beyond a reasonable doubt” that the defendant did it. All of the evidentiary rules are (in theory, at least) designed to ensure that there’s (almost) no chance of wrongfully convicting an innocent defendant. The trial is designed to protect the rights of a defendant: as the English jurist William Blackstone famously said, it is “better that ten guilty persons escape than that one innocent suffer.” In other words, a criminal trial doesn’t treat the two types of error in the same way … punishing the innocent is deemed to be much worse than letting the guilty go free. A statistical test is pretty much the same: the single most important design principle of the test is to *control* the probability of a type I error, to keep it below some fixed probability. This probability, which is denoted α , is called the **significance level** of the test (or sometimes, the *size* of the test). And I’ll say it again, because it is so central to the whole set-up … a hypothesis test is said to have significance level α if the type I error rate is no larger than α .

So, what about the type II error rate? Well, we’d also like to keep those under control too, and we denote this probability by β . However, it’s much more common to refer to the **power** of the test, which is the probability with which we reject a null hypothesis when it really is false, which is $1 - \beta$. To help keep this straight, here’s the same table again, but with the relevant numbers added:

	retain H_0	reject H_0
H_0 is true	$1 - \alpha$ (probability of correct retention)	α (type I error rate)
H_0 is false	β (type II error rate)	$1 - \beta$ (power of the test)

A “powerful” hypothesis test is one that has a small value of β , while still keeping α fixed at some (small) desired level. By convention, scientists make use of three different α levels: .05, .01 and .001. Notice the asymmetry here … the tests are designed to *ensure* that the α level is kept small, but there’s no corresponding guarantee regarding β . We’d certainly *like* the type II error rate to be small, and we try to design tests that keep it small, but this is very much secondary to the overwhelming need to control the type I error rate. As Blackstone might have said if he were a statistician, it is “better to retain 10 false null hypotheses than to reject a single true one”. To be honest, I don’t know that I agree with this philosophy – there are situations where I think it makes sense, and situations where I think it doesn’t – but that’s neither here nor there. It’s how the tests are built.

⁵ An aside regarding the language you use to talk about hypothesis testing. Firstly, one thing you really want to avoid is the word “prove”: a statistical test really doesn’t *prove* that a hypothesis is true or false. Proof implies certainty, and as the saying goes, statistics means never having to say you’re certain. On that point almost everyone would agree. However, beyond that there’s a fair amount of confusion. Some people argue that you’re only allowed to make statements like “rejected the null”, “failed to reject the null”, or possibly “retained the null”. According to this line of thinking, you can’t say things like “accept the alternative” or “accept the null”. Personally I think this is too strong: in my opinion, this conflates null hypothesis testing with Karl Popper’s falsificationist view of the scientific process. While there are similarities between falsificationism and null hypothesis testing, they aren’t equivalent. However, while I personally think it’s fine to talk about accepting a hypothesis (on the proviso that “acceptance” doesn’t actually mean that it’s necessarily true, especially in the case of the null hypothesis), many people will disagree. And more to the point, you should be aware that this particular weirdness exists, so that you’re not caught unawares by it when writing up your own results.

Test statistics and sampling distributions

At this point we need to start talking specifics about how a hypothesis test is constructed. To that end, let's return to the ESP example. Let's ignore the actual data that we obtained, for the moment, and think about the structure of the experiment. Regardless of what the actual numbers are, the *form* of the data is that X out of N people correctly identified the colour of the hidden card. Moreover, let's suppose for the moment that the null hypothesis really is true: ESP doesn't exist, and the true probability that anyone picks the correct colour is exactly $\theta = 0.5$. What would we *expect* the data to look like? Well, obviously, we'd expect the proportion of people who make the correct response to be pretty close to 50%. Or, to phrase this in more mathematical terms, we'd say that X/N is approximately 0.5. Of course, we wouldn't expect this fraction to be *exactly* 0.5: if, for example we tested $N = 100$ people, and $X = 53$ of them got the question right, we'd probably be forced to concede that the data are quite consistent with the null hypothesis. On the other hand, if $X = 99$ of our participants got the question right, then we'd feel pretty confident that the null hypothesis is wrong. Similarly, if only $X = 3$ people got the answer right, we'd be similarly confident that the null was wrong. Let's be a little more technical about this: we have a quantity X that we can calculate by looking at our data; after looking at the value of X , we make a decision about whether to believe that the null hypothesis is correct, or to reject the null hypothesis in favour of the alternative. The name for this thing that we calculate to guide our choices is a **test statistic**.

Having chosen a test statistic, the next step is to state precisely which values of the test statistic would cause us to reject the null hypothesis, and which values would cause us to keep it. In order to do so, we need to determine what the **sampling distribution of the test statistic** would be if the null hypothesis were actually true (we talked about sampling distributions earlier in Section 10.3). Why do we need this? Because this distribution tells us exactly what values of X our null hypothesis would lead us to expect. And therefore, we can use this distribution as a tool for assessing how closely the null hypothesis agrees with our data.

How do we actually determine the sampling distribution of the test statistic? For a lot of hypothesis tests this step is actually quite complicated, and later on in the book you'll see me being slightly evasive about it for some of the tests (some of them I don't even understand myself). However, sometimes it's very easy. And, fortunately for us, our ESP example provides us with one of the easiest cases. Our population parameter θ is just the overall probability that people respond correctly when asked the question, and our test statistic X is the *count* of the number of people who did so, out of a sample size of N . We've seen a distribution like this before, in Section 9.3: that's exactly what the binomial distribution describes! So, to use the notation and terminology that I introduced in that section, we would say that the null hypothesis predicts that X is binomially distributed, which is written

$$X \sim \text{Binomial}(\theta, N)$$

Since the null hypothesis states that $\theta = 0.5$ and our experiment has $N = 100$ people, we have the sampling distribution we need. This sampling distribution is plotted in Figure 11.1. No surprises really: the null hypothesis says that $X = 50$ is the most likely outcome, and it says that we're almost certain to see somewhere between 40 and 60 correct responses.

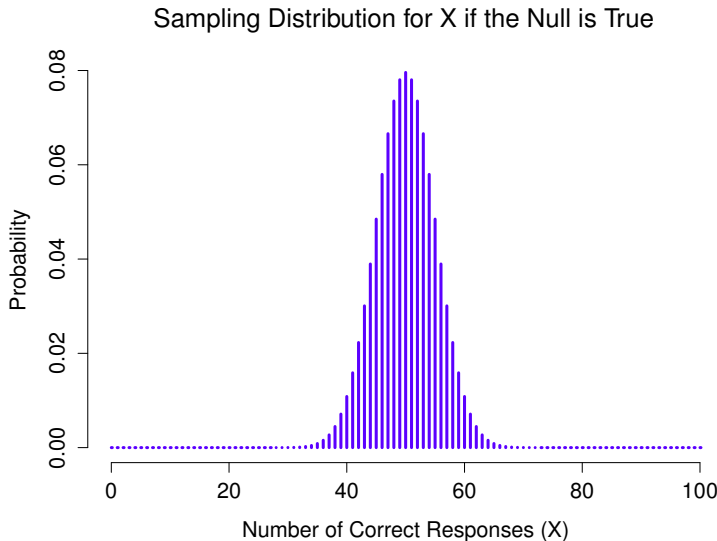


Figure 11.1: The sampling distribution for our test statistic X when the null hypothesis is true. For our ESP scenario, this is a binomial distribution. Not surprisingly, since the null hypothesis says that the probability of a correct response is $\theta = .5$, the sampling distribution says that the most likely value is 50 (out of 100) correct responses. Most of the probability mass lies between 40 and 60.

11.4 _____

Making decisions

Okay, we're very close to being finished. We've constructed a test statistic (X), and we chose this test statistic in such a way that we're pretty confident that if X is close to $N/2$ then we should retain the null, and if not we should reject it. The question that remains is this: exactly which values of the test statistic should we associate with the null hypothesis, and which exactly values go with the alternative hypothesis? In my ESP study, for example, I've observed a value of $X = 62$. What decision should I make? Should I choose to believe the null hypothesis, or the alternative hypothesis?

11.4.1 Critical regions and critical values

To answer this question, we need to introduce the concept of a **critical region** for the test statistic X . The critical region of the test corresponds to those values of X that would lead us to reject null hypothesis (which is why the critical region is also sometimes called the rejection region). How do we find this critical region? Well, let's consider what we know:

1. X should be very big or very small in order to reject the null hypothesis.
2. If the null hypothesis is true, the sampling distribution of X is $\text{Binomial}(0.5, N)$.
3. If $\alpha = .05$, the critical region must cover 5% of this sampling distribution.

Critical Regions for a Two-Sided Test

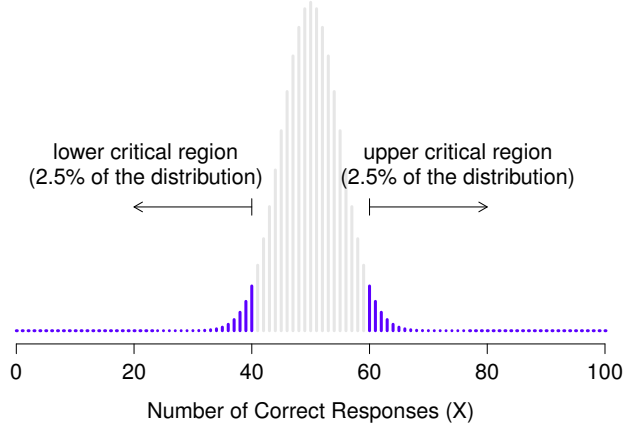


Figure 11.2: The critical region associated with the hypothesis test for the ESP study, for a hypothesis test with a significance level of $\alpha = .05$. The plot itself shows the sampling distribution of X under the null hypothesis (i.e., same as Figure 11.1): the grey bars correspond to those values of X for which we would retain the null hypothesis. The black bars show the critical region: those values of X for which we would reject the null. Because the alternative hypothesis is two sided (i.e., allows both $\theta < .5$ and $\theta > .5$), the critical region covers both tails of the distribution. To ensure an α level of $.05$, we need to ensure that each of the two regions encompasses 2.5% of the sampling distribution.

It's important to make sure you understand this last point: the critical region corresponds to those values of X for which we would reject the null hypothesis, and the sampling distribution in question describes the probability that we would obtain a particular value of X if the null hypothesis were actually true. Now, let's suppose that we chose a critical region that covers 20% of the sampling distribution, and suppose that the null hypothesis is actually true. What would be the probability of incorrectly rejecting the null? The answer is of course 20% . And therefore, we would have built a test that had an α level of 0.2 . If we want $\alpha = .05$, the critical region is only *allowed* to cover 5% of the sampling distribution of our test statistic.

As it turns out, those three things uniquely solve the problem: our critical region consists of the most *extreme values*, known as the **tails** of the distribution. This is illustrated in Figure 11.2. As it turns out, if we want $\alpha = .05$, then our critical regions correspond to $X \leq 40$ and $X \geq 60$.⁶ That is, if the number of people saying "true" is between 41 and 59, then we should retain the null hypothesis. If the number is between 0 to 40 or between 60 to 100, then we should reject the null hypothesis. The numbers 40 and 60 are often referred to as the **critical values**, since they define the edges of the critical region.

At this point, our hypothesis test is essentially complete: (1) we choose an α level (e.g., $\alpha = .05$), (2) come up with some test statistic (e.g., X) that does a good job (in some meaningful sense) of comparing H_0 to H_1 , (3) figure out the sampling distribution of the test statistic on the assumption that the null hypothesis is true (in this case, binomial) and then (4) calculate the critical region that produces an

⁶Strictly speaking, the test I just constructed has $\alpha = .057$, which is a bit too generous. However, if I'd chosen 39 and 61 to be the boundaries for the critical region, then the critical region only covers 3.5% of the distribution. I figured that it makes more sense to use 40 and 60 as my critical values, and be willing to tolerate a 5.7% type I error rate, since that's as close as I can get to a value of $\alpha = .05$.

appropriate α level (0-40 and 60-100). All that we have to do now is calculate the value of the test statistic for the real data (e.g., $X = 62$) and then compare it to the critical values to make our decision. Since 62 is greater than the critical value of 60, we would reject the null hypothesis. Or, to phrase it slightly differently, we say that the test has produced a **significant** result.

11.4.2 A note on statistical “significance”

Like other occult techniques of divination, the statistical method has a private jargon deliberately contrived to obscure its methods from non-practitioners.

– G. O. Ashley

A very brief digression is in order at this point, regarding the word “significant”. The concept of statistical significance is actually a very simple one, but has a very unfortunate name. If the data allow us to reject the null hypothesis, we say that “the result is *statistically significant*”, which is often shortened to “the result is significant”. This terminology is rather old, and dates back to a time when “significant” just meant something like “indicated”, rather than its modern meaning, which is much closer to “important”. As a result, a lot of modern readers get very confused when they start learning statistics, because they think that a “significant result” must be an important one. It doesn’t mean that at all. All that “statistically significant” means is that the data allowed us to reject a null hypothesis. Whether or not the result is actually important in the real world is a very different question, and depends on all sorts of other things.

11.4.3 The difference between one sided and two sided tests

There’s one more thing I want to point out about the hypothesis test that I’ve just constructed. If we take a moment to think about the statistical hypotheses I’ve been using,

$$\begin{aligned} H_0 : \theta &= .5 \\ H_1 : \theta &\neq .5 \end{aligned}$$

we notice that the alternative hypothesis covers *both* the possibility that $\theta < .5$ and the possibility that $\theta > .5$. This makes sense if I really think that ESP could produce better-than-chance performance *or* worse-than-chance performance (and there are some people who think that). In statistical language, this is an example of a **two-sided test**. It’s called this because the alternative hypothesis covers the area on both “sides” of the null hypothesis, and as a consequence the critical region of the test covers both tails of the sampling distribution (2.5% on either side if $\alpha = .05$), as illustrated earlier in Figure 11.2.

However, that’s not the only possibility. It might be the case, for example, that I’m only willing to believe in ESP if it produces better than chance performance. If so, then my alternative hypothesis would only covers the possibility that $\theta > .5$, and as a consequence the null hypothesis now becomes $\theta \leq .5$:

$$\begin{aligned} H_0 : \theta &\leq .5 \\ H_1 : \theta &> .5 \end{aligned}$$

Critical Region for a One-Sided Test

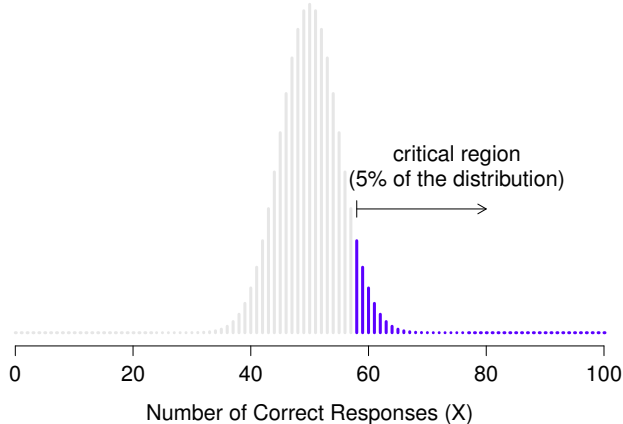


Figure 11.3: The critical region for a one sided test. In this case, the alternative hypothesis is that $\theta > .05$, so we would only reject the null hypothesis for large values of X . As a consequence, the critical region only covers the upper tail of the sampling distribution; specifically the upper 5% of the distribution. Contrast this to the two-sided version in Figure 11.2.

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When this happens, we have what's called a **one-sided test**, and when this happens the critical region only covers one tail of the sampling distribution. This is illustrated in Figure 11.3.

11.5

The p value of a test

In one sense, our hypothesis test is complete; we've constructed a test statistic, figured out its sampling distribution if the null hypothesis is true, and then constructed the critical region for the test. Nevertheless, I've actually omitted the most important number of all: **the p value**. It is to this topic that we now turn. There are two somewhat different ways of interpreting a p value, one proposed by Sir Ronald Fisher and the other by Jerzy Neyman. Both versions are legitimate, though they reflect very different ways of thinking about hypothesis tests. Most introductory textbooks tend to give Fisher's version only, but I think that's a bit of a shame. To my mind, Neyman's version is cleaner, and actually better reflects the logic of the null hypothesis test. You might disagree though, so I've included both. I'll start with Neyman's version...

11.5.1 A softer view of decision making

One problem with the hypothesis testing procedure that I've described is that it makes no distinction at all between a result this “barely significant” and those that are “highly significant”. For instance, in my ESP study the data I obtained only just fell inside the critical region - so I did get a significant effect, but was a pretty near thing. In contrast, suppose that I'd run a study in which $X = 97$ out of my $N = 100$ participants got the answer right. This would obviously be significant too, but my a much

larger margin; there's really no ambiguity about this at all. The procedure that I described makes no distinction between the two. If I adopt the standard convention of allowing $\alpha = .05$ as my acceptable Type I error rate, then both of these are significant results.

This is where the p value comes in handy. To understand how it works, let's suppose that we ran lots of hypothesis tests on the same data set: but with a different value of α in each case. When we do that for my original ESP data, what we'd get is something like this

Value of α	0.05	0.04	0.03	0.02	0.01
Reject the null?	Yes	Yes	Yes	No	No

When we test ESP data ($X = 62$ successes out of $N = 100$ observations) using α levels of .03 and above, we'd always find ourselves rejecting the null hypothesis. For α levels of .02 and below, we always end up retaining the null hypothesis. Therefore, somewhere between .02 and .03 there must be a smallest value of α that would allow us to reject the null hypothesis for this data. This is the p value; as it turns out the ESP data has $p = .021$. In short:

p is defined to be the smallest Type I error rate (α) that you have to be willing to tolerate if you want to reject the null hypothesis.

If it turns out that p describes an error rate that you find intolerable, then you must retain the null. If you're comfortable with an error rate equal to p , then it's okay to reject the null hypothesis in favour of your preferred alternative.

In effect, p is a summary of all the possible hypothesis tests that you could have run, taken across all possible α values. And as a consequence it has the effect of “softening” our decision process. For those tests in which $p \leq \alpha$ you would have rejected the null hypothesis, whereas for those tests in which $p > \alpha$ you would have retained the null. In my ESP study I obtained $X = 62$, and as a consequence I’ve ended up with $p = .021$. So the error rate I have to tolerate is 2.1%. In contrast, suppose my experiment had yielded $X = 97$. What happens to my p value now? This time it’s shrunk to $p = 1.36 \times 10^{-25}$, which is a tiny, tiny⁷ Type I error rate. For this second case I would be able to reject the null hypothesis with a lot more confidence, because I only have to be “willing” to tolerate a type I error rate of about 1 in 10 trillion trillion in order to justify my decision to reject.

11.5.2 The probability of extreme data

The second definition of the p -value comes from Sir Ronald Fisher, and it's actually this one that you tend to see in most introductory statistics textbooks. Notice how, when I constructed the critical region, it corresponded to the *tails* (i.e., extreme values) of the sampling distribution? That's not a coincidence: almost all "good" tests have this characteristic (good in the sense of minimising our type II error rate, β). The reason for that is that a good critical region almost always corresponds to those values of the test statistic that are least likely to be observed if the null hypothesis is true. If this rule is true, then we can define the p -value as the probability that we would have observed a test statistic that is at least as extreme as the one we actually did get. In other words, if the data are extremely implausible

according to the null hypothesis, then the null hypothesis is probably wrong.⁸

11.5.3 A common mistake

Okay, so you can see that there are two rather different but legitimate ways to interpret the p value, one based on Neyman's approach to hypothesis testing and the other based on Fisher's. Unfortunately, there is a third explanation that people sometimes give, especially when they're first learning statistics, and it is *absolutely and completely wrong*. This mistaken approach is to refer to the p value as "the probability that the null hypothesis is true". It's an intuitively appealing way to think, but it's wrong in two key respects: (1) null hypothesis testing is a frequentist tool, and the frequentist approach to probability does *not* allow you to assign probabilities to the null hypothesis... according to this view of probability, the null hypothesis is either true or it is not; it cannot have a "5% chance" of being true. (2) even within the Bayesian approach, which does let you assign probabilities to hypotheses, the p value would not correspond to the probability that the null is true; this interpretation is entirely inconsistent with the mathematics of how the p value is calculated. Put bluntly, despite the intuitive appeal of thinking this way, there is no justification for interpreting a p value this way. Never do it.

11.6

Reporting the results of a hypothesis test

When writing up the results of a hypothesis test, there's usually several pieces of information that you need to report, but it varies a fair bit from test to test. Throughout the rest of the book I'll spend a little time talking about how to report the results of different tests (see Section 12.1.8 for a particularly detailed example), so that you can get a feel for how it's usually done. However, regardless of what test you're doing, the one thing that you always have to do is say something about the p value, and whether or not the outcome was significant.

The fact that you have to do this is unsurprising; it's the whole point of doing the test. What might be surprising is the fact that there is some contention over exactly how you're supposed to do it. Leaving aside those people who completely disagree with the entire framework underpinning null hypothesis testing, there's a certain amount of tension that exists regarding whether or not to report the exact p value that you obtained, or if you should state only that $p < \alpha$ for a significance level that you chose in advance (e.g., $p < .05$).

11.6.1 The issue

To see why this is an issue, the key thing to recognise is that p values are *terribly* convenient. In practice, the fact that we can compute a p value means that we don't actually have to specify any α level at all in order to run the test. Instead, what you can do is calculate your p value and interpret it directly: if you get $p = .062$, then it means that you'd have to be willing to tolerate a Type I error rate of 6.2% to justify rejecting the null. If you personally find 6.2% intolerable, then you retain the null. Therefore, the argument goes, why don't we just report the actual p value and let the reader make up their own minds about what an acceptable Type I error rate is? This approach has the big advantage of "softening" the decision making process – in fact, if you accept the Neyman definition of the p value, that's the whole point of the p value. We no longer have a fixed significance level of $\alpha = .05$ as a bright line separating

⁸You might wonder why I don't like Fisher's version as much as Neyman's. Basically, it's because Fisher's definition only works for 99.99% of the null hypothesis tests that I can think of, while Neyman's version works for 100% of cases. There are a few pathological ways of choosing your null hypothesis and alternative hypothesis in such a way that Fisher's definition breaks. I won't go into them here, other than to say that it is possible.

“accept” from “reject” decisions; and this removes the rather pathological problem of being forced to treat $p = .051$ in a fundamentally different way to $p = .049$.

This flexibility is both the advantage and the disadvantage to the p value. The reason why a lot of people don’t like the idea of reporting an exact p value is that it gives the researcher a bit *too much* freedom. In particular, it lets you change your mind about what error tolerance you’re willing to put up with *after* you look at the data. For instance, consider my ESP experiment. Suppose I ran my test, and ended up with a p value of .09. Should I accept or reject? Now, to be honest, I haven’t yet bothered to think about what level of Type I error I’m “really” willing to accept. I don’t have an opinion on that topic. But I *do* have an opinion about whether or not ESP exists, and I *definitely* have an opinion about whether my research should be published in a reputable scientific journal. And amazingly, now that I’ve looked at the data I’m starting to think that a 9% error rate isn’t so bad, especially when compared to how annoying it would be to have to admit to the world that my experiment has failed. So, to avoid looking like I just made it up after the fact, I now say that my α is .1: a 10% type I error rate isn’t too bad, and at that level my test is significant! I win.

In other words, the worry here is that I might have the best of intentions, and be the most honest of people, but the temptation to just “shade” things a little bit here and there is really, really strong. As anyone who has ever run an experiment can attest, it’s a long and difficult process, and you often get *very* attached to your hypotheses. It’s hard to let go and admit the experiment didn’t find what you wanted it to find. And that’s the danger here. If we use the “raw” p -value, people will start interpreting the data in terms of what they *want* to believe, not what the data are actually saying... and if we allow that, well, why are we bothering to do science at all? Why not let everyone believe whatever they like about anything, regardless of what the facts are? Okay, that’s a bit extreme, but that’s where the worry comes from. According to this view, you really *must* specify your α value in advance, and then only report whether the test was significant or not. It’s the only way to keep ourselves honest.

11.6.2 Two proposed solutions

In practice, it’s pretty rare for a researcher to specify a single α level ahead of time. Instead, the convention is that scientists rely on three standard significance levels: .05, .01 and .001. When reporting your results, you indicate which (if any) of these significance levels allow you to reject the null hypothesis. This is summarised in Table 11.1. This allows us to soften the decision rule a little bit, since $p < .01$ implies that the data meet a stronger evidentiary standard than $p < .05$ would. Nevertheless, since these levels are fixed in advance by convention, it does prevent people choosing their α level after looking at the data.

Nevertheless, quite a lot of people still prefer to report exact p values. To many people, the advantage of allowing the reader to make up their own mind about how to interpret $p = .06$ outweighs any disadvantages. In practice, however, even among those researchers who prefer exact p values it is quite common to just write $p < .001$ instead of reporting an exact value for small p . This is in part because a lot of software doesn’t actually print out the p value when it’s that small (e.g., SPSS just writes $p = .000$ whenever $p < .001$), and in part because a very small p value can be kind of misleading. The human mind sees a number like .000000001 and it’s hard to suppress the gut feeling that the evidence in favour of the alternative hypothesis is a near certainty. In practice however, this is usually wrong. Life is a big, messy, complicated thing; and every statistical test ever invented relies on simplifications, approximations and assumptions. As a consequence, it’s probably not reasonable to walk away from *any* statistical analysis with a feeling of confidence stronger than $p < .001$ implies. In other words, $p < .001$ is really code for “as far as *this test* is concerned, the evidence is overwhelming.”

In light of all this, you might be wondering exactly what you should do. There’s a fair bit of contradictory advice on the topic, with some people arguing that you should report the exact p value, and other people arguing that you should use the tiered approach illustrated in Table 11.1. As a result, the

Table 11.1: A commonly adopted convention for reporting p values: in many places it is conventional to report one of four different things (e.g., $p < .05$) as shown below. I've included the “significance stars” notation (i.e., a * indicates $p < .05$) because you sometimes see this notation produced by statistical software. It's also worth noting that some people will write *n.s.* (not significant) rather than $p > .05$.

Usual notation	Signif. stars	English translation	The null is...
$p > .05$		The test wasn't significant	Retained
$p < .05$	*	The test was significant at $\alpha = .05$ but not at $\alpha = .01$ or $\alpha = .001$.	Rejected
$p < .01$	**	The test was significant at $\alpha = .05$ and $\alpha = .01$ but not at $\alpha = .001$.	Rejected
$p < .001$	***	The test was significant at all levels	Rejected
.....			

best advice I can give is to suggest that you look at papers/reports written in your field and see what the convention seems to be. If there doesn't seem to be any consistent pattern, then use whichever method you prefer.

11.7

Running the hypothesis test in practice

At this point some of you might be wondering if this is a “real” hypothesis test, or just a toy example that I made up. It’s real. In the previous discussion I built the test from first principles, thinking that it was the simplest possible problem that you might ever encounter in real life. However, this test already exists: it’s called the *binomial test*, and it’s implemented by an R function called `binom.test()`. To test the null hypothesis that the response probability is one-half $p = .5$,⁹ using data in which $x = 62$ of $n = 100$ people made the correct response, here’s how to do it in R:

```
> binom.test( x=62, n=100, p=.5 )

Exact binomial test

data: 62 and 100
number of successes = 62, number of trials = 100,
p-value = 0.02098
alternative hypothesis: true probability of success is not equal to 0.5
95 percent confidence interval:
 0.5174607 0.7152325
sample estimates:
probability of success
                0.62
```

Right now, this output looks pretty unfamiliar to you, but you can see that it’s telling you more or less the right things. Specifically, the p -value of 0.02 is less than the usual choice of $\alpha = .05$, so you can reject

⁹Note that the `p` here has nothing to do with a p value. The `p` argument in the `binom.test()` function corresponds to the probability of making a correct response, according to the null hypothesis. In other words, it’s the θ value.

Sampling Distribution for X if $\theta = .55$

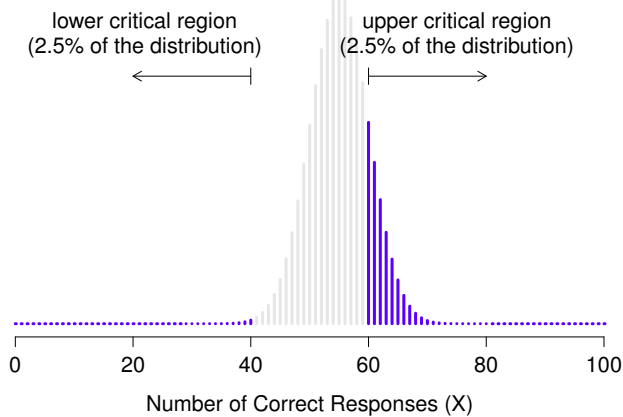


Figure 11.4: Sampling distribution under the *alternative* hypothesis, for a population parameter value of $\theta = 0.55$. A reasonable proportion of the distribution lies in the rejection region.

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the null. We'll talk a lot more about how to read this sort of output as we go along; and after a while you'll hopefully find it quite easy to read and understand. For now, however, I just wanted to make the point that R contains a whole lot of functions corresponding to different kinds of hypothesis test. And while I'll usually spend quite a lot of time explaining the logic behind how the tests are built, every time I discuss a hypothesis test the discussion will end with me showing you a fairly simple R command that you can use to run the test in practice.

11.8 _____

Effect size, sample size and power

In previous sections I've emphasised the fact that the major design principle behind statistical hypothesis testing is that we try to control our Type I error rate. When we fix $\alpha = .05$ we are attempting to ensure that only 5% of true null hypotheses are incorrectly rejected. However, this doesn't mean that we don't care about Type II errors. In fact, from the researcher's perspective, the error of failing to reject the null when it is actually false is an extremely annoying one. With that in mind, a secondary goal of hypothesis testing is to try to minimise β , the Type II error rate, although we don't usually *talk* in terms of minimising Type II errors. Instead, we talk about maximising the *power* of the test. Since power is defined as $1 - \beta$, this is the same thing.

11.8.1 The power function

Let's take a moment to think about what a Type II error actually is. A Type II error occurs when the alternative hypothesis is true, but we are nevertheless unable to reject the null hypothesis. Ideally, we'd be able to calculate a single number β that tells us the Type II error rate, in the same way that we can

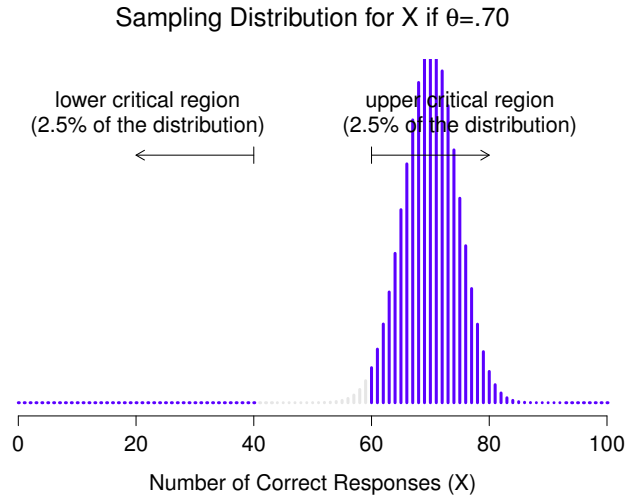


Figure 11.5: Sampling distribution under the *alternative* hypothesis, for a population parameter value of $\theta = 0.70$. Almost all of the distribution lies in the rejection region.

set $\alpha = .05$ for the Type I error rate. Unfortunately, this is a lot trickier to do. To see this, notice that in my ESP study the alternative hypothesis actually corresponds to lots of possible values of θ . In fact, the alternative hypothesis corresponds to every value of θ *except* 0.5. Let's suppose that the true probability of someone choosing the correct response is 55% (i.e., $\theta = .55$). If so, then the *true* sampling distribution for X is not the same one that the null hypothesis predicts: the most likely value for X is now 55 out of 100. Not only that, the whole sampling distribution has now shifted, as shown in Figure 11.4. The critical regions, of course, do not change: by definition, the critical regions are based on what the null hypothesis predicts. What we're seeing in this figure is the fact that when the null hypothesis is wrong, a much larger proportion of the sampling distribution falls in the critical region. And of course that's what should happen: the probability of rejecting the null hypothesis is larger when the null hypothesis is actually false! However $\theta = .55$ is not the only possibility consistent with the alternative hypothesis. Let's instead suppose that the true value of θ is actually 0.7. What happens to the sampling distribution when this occurs? The answer, shown in Figure 11.5, is that almost the entirety of the sampling distribution has now moved into the critical region. Therefore, if $\theta = 0.7$ the probability of us correctly rejecting the null hypothesis (i.e., the power of the test) is much larger than if $\theta = 0.55$. In short, while $\theta = .55$ and $\theta = .70$ are both part of the alternative hypothesis, the Type II error rate is different.

What all this means is that the power of a test (i.e., $1 - \beta$) depends on the true value of θ . To illustrate this, I've calculated the expected probability of rejecting the null hypothesis for all values of θ , and plotted it in Figure 11.6. This plot describes what is usually called the **power function** of the test. It's a nice summary of how good the test is, because it actually tells you the power ($1 - \beta$) for all possible values of θ . As you can see, when the true value of θ is very close to 0.5, the power of the test drops very sharply, but when it is further away, the power is large.

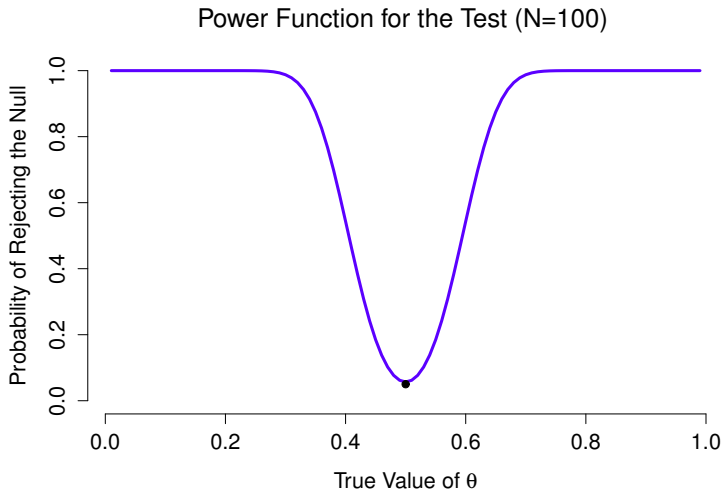


Figure 11.6: The probability that we will reject the null hypothesis, plotted as a function of the true value of θ . Obviously, the test is more powerful (greater chance of correct rejection) if the true value of θ is very different from the value that the null hypothesis specifies (i.e., $\theta = .5$). Notice that when θ actually is equal to .05 (plotted as a black dot), the null hypothesis is in fact true: rejecting the null hypothesis in this instance would be a Type I error.

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11.8.2 Effect size

It is inappropriate to be concerned with mice when there are tigers abroad
– George Box

The plot shown in Figure 11.6 captures a fairly basic point about hypothesis testing. If the true state of the world is very different from what the null hypothesis predicts, then your power will be very high; but if the true state of the world is similar to the null (but not identical) then the power of the test is going to be very low. Therefore, it's useful to be able to have some way of quantifying how "similar" the true state of the world is to the null hypothesis. A statistic that does this is called a measure of **effect size** (e.g., Cohen, 1988; Ellis, 2010). Effect size is defined slightly differently in different contexts,¹⁰ (and so this section just talks in general terms) but the qualitative idea that it tries to capture is always the same: how big is the difference between the *true* population parameters, and the parameter values that are assumed by the null hypothesis? In our ESP example, if we let $\theta_0 = 0.5$ denote the value assumed by the null hypothesis, and let θ denote the true value, then a simple measure of effect size could be something like the difference between the true value and null (i.e., $\theta - \theta_0$), or possibly just the magnitude of this difference, $\text{abs}(\theta - \theta_0)$.

Why calculate effect size? Let's assume that you've run your experiment, collected the data, and gotten a significant effect when you ran your hypothesis test. Isn't it enough just to say that you've gotten a significant effect? Surely that's the *point* of hypothesis testing? Well, sort of. Yes, the point of doing a hypothesis test is to try to demonstrate that the null hypothesis is wrong, but that's hardly the

¹⁰There's an R package called `compute.es` that can be used for calculating a very broad range of effect size measures; but for the purposes of the current book we won't need it: all of the effect size measures that I'll talk about here have functions in the `lsr` package

Table 11.2: A crude guide to understanding the relationship between statistical significance and effect sizes. Basically, if you don't have a significant result, then the effect size is pretty meaningless; because you don't have any evidence that it's even real. On the other hand, if you do have a significant effect but your effect size is small, then there's a pretty good chance that your result (although real) isn't all that interesting. However, this guide is very crude: it depends a lot on what exactly you're studying. Small effects can be of massive practical importance in some situations. So don't take this table too seriously. It's a rough guide at best.

	big effect size	small effect size
significant result	difference is real, and of practical importance	difference is real, but might not be interesting
non-significant result	no effect observed	no effect observed

only thing we're interested in. If the null hypothesis claimed that $\theta = .5$, and we show that it's wrong, we've only really told half of the story. Rejecting the null hypothesis implies that we believe that $\theta \neq .5$, but there's a big difference between $\theta = .51$ and $\theta = .8$. If we find that $\theta = .8$, then not only have we found that the null hypothesis is wrong, it appears to be *very* wrong. On the other hand, suppose we've successfully rejected the null hypothesis, but it looks like the true value of θ is only $.51$ (this would only be possible with a large study). Sure, the null hypothesis is wrong, but it's not at all clear that we actually *care*, because the effect size is so small. In the context of my ESP study we might still care, since any demonstration of real psychic powers would actually be pretty cool¹¹, but in other contexts a 1% difference isn't very interesting, even if it is a real difference. For instance, suppose we're looking at differences in high school exam scores between males and females, and it turns out that the female scores are 1% higher on average than the males. If I've got data from thousands of students, then this difference will almost certainly be *statistically significant*, but regardless of how small the p value is it's just not very interesting. You'd hardly want to go around proclaiming a crisis in boys education on the basis of such a tiny difference would you? It's for this reason that it is becoming more standard (slowly, but surely) to report some kind of standard measure of effect size along with the results of the hypothesis test. The hypothesis test itself tells you whether you should believe that the effect you have observed is real (i.e., not just due to chance); the effect size tells you whether or not you should care.

11.8.3 Increasing the power of your study

Not surprisingly, scientists are fairly obsessed with maximising the power of their experiments. We want our experiments to work, and so we want to maximise the chance of rejecting the null hypothesis if it is false (and of course we usually want to believe that it is false!) As we've seen, one factor that influences power is the effect size. So the first thing you can do to increase your power is to increase the effect size. In practice, what this means is that you want to design your study in such a way that the effect size gets magnified. For instance, in my ESP study I might believe that psychic powers work best in a quiet, darkened room; with fewer distractions to cloud the mind. Therefore I would try to conduct my experiments in just such an environment: if I can strengthen people's ESP abilities somehow, then

¹¹Although in practice a very small effect size is worrying, because even very minor methodological flaws might be responsible for the effect; and in practice no experiment is perfect, so there are always methodological issues to worry about.

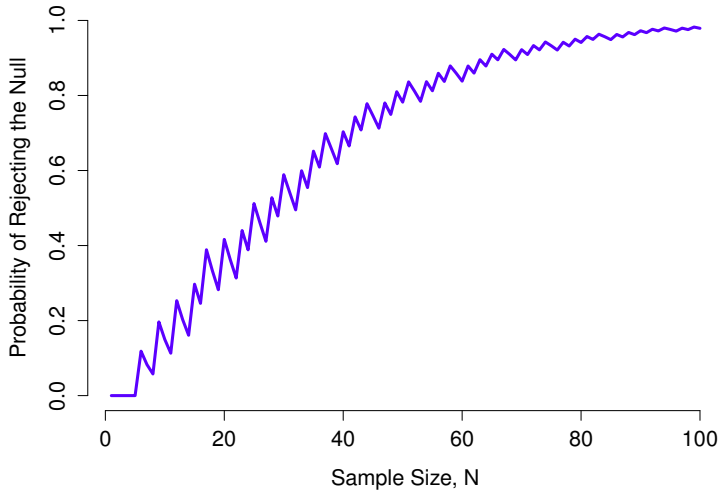


Figure 11.7: The power of our test, plotted as a function of the sample size N . In this case, the true value of θ is 0.7, but the null hypothesis is that $\theta = 0.5$. Overall, larger N means greater power. (The small zig-zags in this function occur because of some odd interactions between θ , α and the fact that the binomial distribution is discrete; it doesn't matter for any serious purpose)

the true value of θ will go up¹² and therefore my effect size will be larger. In short, clever experimental design is one way to boost power; because it can alter the effect size.

Unfortunately, it's often the case that even with the best of experimental designs you may have only a small effect. Perhaps, for example, ESP really does exist, but even under the best of conditions it's very very weak. Under those circumstances, your best bet for increasing power is to increase the sample size. In general, the more observations that you have available, the more likely it is that you can discriminate between two hypotheses. If I ran my ESP experiment with 10 participants, and 7 of them correctly guessed the colour of the hidden card, you wouldn't be terribly impressed. But if I ran it with 10,000 participants and 7,000 of them got the answer right, you would be much more likely to think I had discovered something. In other words, power increases with the sample size. This is illustrated in Figure 11.7, which shows the power of the test for a true parameter of $\theta = 0.7$, for all sample sizes N from 1 to 100, where I'm assuming that the null hypothesis predicts that $\theta_0 = 0.5$.

Because power is important, whenever you're contemplating running an experiment it would be pretty useful to know how much power you're likely to have. It's never possible to know for sure, since you can't possibly know what your effect size is. However, it's often (well, sometimes) possible to guess how big it should be. If so, you can guess what sample size you need! This idea is called **power analysis**, and if it's feasible to do it, then it's very helpful, since it can tell you something about whether you have enough time or money to be able to run the experiment successfully. It's increasingly common to see people arguing that power analysis should be a required part of experimental design, so it's worth knowing about. I don't discuss power analysis in this book, however. This is partly for a boring reason

¹²Notice that the true population parameter θ doesn't necessarily correspond to an immutable fact of nature. In this context θ is just the true probability that people would correctly guess the colour of the card in the other room. As such the population parameter can be influenced by all sorts of things. Of course, this is all on the assumption that ESP actually exists!

and partly for a substantive one. The boring reason is that I haven't had time to write about power analysis yet. The substantive one is that I'm still a little suspicious of power analysis. Speaking as a researcher, I have very rarely found myself in a position to be able to do one – it's either the case that (a) my experiment is a bit non-standard and I don't know how to define effect size properly, (b) I literally have so little idea about what the effect size will be that I wouldn't know how to interpret the answers. Not only that, after extensive conversations with someone who does stats consulting for a living (my wife, as it happens), I can't help but notice that in practice the *only* time anyone ever asks her for a power analysis is when she's helping someone write a grant application. In other words, the only time any scientist ever seems to want a power analysis in real life is when they're being forced to do it by bureaucratic process. It's not part of anyone's day to day work. In short, I've always been of the view that while power is an important concept, power *analysis* is not as useful as people make it sound, except in the rare cases where (a) someone has figured out how to calculate power for your actual experimental design and (b) you have a pretty good idea what the effect size is likely to be. Maybe other people have had better experiences than me, but I've personally never been in a situation where both (a) and (b) were true. Maybe I'll be convinced otherwise in the future, and probably a future version of this book would include a more detailed discussion of power analysis, but for now this is about as much as I'm comfortable saying about the topic.

11.9

Some issues to consider

What I've described to you in this chapter is the orthodox framework for null hypothesis significance testing (NHST). Understanding how NHST works is an absolute necessity, since it has been the dominant approach to inferential statistics ever since it came to prominence in the early 20th century. It's what the vast majority of working scientists rely on for their data analysis, so even if you hate it you need to know it. However, the approach is not without problems. There are a number of quirks in the framework, historical oddities in how it came to be, theoretical disputes over whether or not the framework is right, and a lot of practical traps for the unwary. I'm not going to go into a lot of detail on this topic, but I think it's worth briefly discussing a few of these issues.

11.9.1 Neyman versus Fisher

The first thing you should be aware of is that orthodox NHST is actually a mash-up of two rather different approaches to hypothesis testing, one proposed by Sir Ronald Fisher and the other proposed by Jerzy Neyman. Here's a quick summary of what I take these two approaches to be. First, let's talk about Fisher's approach. As far as I can tell, Fisher assumed that you only had the one hypothesis (the null), and what you want to do is find out if the null hypothesis is inconsistent with the data. From his perspective, what you should do is check to see if the data are "sufficiently unlikely" according to the null. In fact, if you remember back to our earlier discussion, that's how Fisher defines the *p*-value. According to Fisher, if the null hypothesis provided a very poor account of the data, you could safely reject it. But, since you don't have any other hypotheses to compare it to, there's no way of "accepting the alternative" because you don't necessarily have an explicitly stated alternative. That's more or less all that there was to it.

In contrast, Neyman thought that the point of hypothesis testing was as a guide to action, and his approach was somewhat more formal than Fisher's. His view was that there are multiple things that you could *do* (accept the null or accept the alternative) and the point of the test was to tell you which one the data support. From this perspective, it is critical to specify your alternative hypothesis properly. If you don't know what the alternative hypothesis is, then you don't know how powerful the test is, or even

which action makes sense. His framework genuinely requires a competition between different hypotheses. For Neyman, the p value didn't directly measure the probability of the data (or data more extreme) under the null, it was more of an abstract description about which “possible tests” were telling you to accept the null, and which “possible tests” were telling you to accept the alternative.

As you can see, what we have today is an odd mishmash of the two. We talk about having both a null hypothesis and an alternative (Neyman), but usually¹³ define the p value in terms of extreme data (Fisher), but we still have α values (Neyman). Some of the statistical tests have explicitly specified alternatives (Neyman) but others are quite vague about it (Fisher). And, according to some people at least, we're not allowed to talk about accepting the alternative (Fisher). It's a mess: but I hope this at least explains why it's a mess.

11.9.2 Bayesians versus frequentists

Earlier on in this chapter I was quite emphatic about the fact that you *cannot* interpret the p value as the probability that the null hypothesis is true. NHST is fundamentally a frequentist tool (see Chapter 9) and as such it does not allow you to assign probabilities to hypotheses: the null hypothesis is either true or it is not. The Bayesian approach to statistics interprets probability as a degree of belief, so it's totally okay to say that there is a 10% chance that the null hypothesis is true: that's just a reflection of the degree of confidence that you have in this hypothesis. In modern times, the Bayesian view is an entirely respectable way to do statistics, but it never used to be. Advocates of the Bayesian view tend to point out that there are a lot of undesirable behaviours in the NHST framework; detractors point to undesirable flaws in the Bayesian framework. While I personally tend to side with the Bayesians, I'm not going to launch into a full discussion of the dispute. Right now, all I want to do is point out that this disagreement exists, and to give you a brief flavour of how the Bayesian approach works. If you are interested in this topic, a good place to start is the book by Kruschke (2011).

The Bayesian approach starts from the basic rule of probability theory that states that the joint probability of A happening and B happening, $P(A, B)$, is equal to the probability of one event happening $P(A)$ multiplied by the conditional probability of the other event happening $P(B|A)$ given that we know that the first one has happened (or will happen - the timing of events is actually irrelevant to the maths, so long as we have the relevant knowledge).¹⁴ From this basic rule, we can derive a slightly different way of thinking about this, known as **Bayes' theorem**:

$$P(A|B) = \frac{P(B|A)P(A)}{P(B)}$$

When we talk about it in terms of abstract events A and B , everyone agrees that it is a completely valid rule of probability theory. However, if we let “event B ” refer to “we observe the data X ” and let “event A ” refer to the event that “the null hypothesis H_0 is true”, it sparks a lot of disagreement. Why? Because it means we have to assign *probabilities* to the event that “the null hypothesis is true”. Remember, if you're a frequentist, a probability can only be defined in terms of what happens after a large number of independent replications (i.e., a long run frequency). If this is your interpretation of probability, talking about the “probability” that the null hypothesis is true is complete gibberish: a null hypothesis is either true or it is false. There's no way you can talk about a long run frequency for this statement. To talk

¹³Although this book describes both Neyman's and Fisher's definition of the p value, most don't. Most introductory textbooks will only give you the Fisher version.

¹⁴Back in Chapter 9 I used the notation $P(A \cap B)$ to refer to the probability that both A and B occur, whereas in this chapter I've used $P(A, B)$. This is intentional. In probability theory it's traditional to use $P(A \cap B)$, but in statistics it's more common to shorten it to $P(A, B)$. So the chapter on probability theory uses the standard notation from probability theory, whereas the chapters on statistics tend to use the statistical notation. In an ideal world the notation would be the same, but I think it's pretty clear that we don't live in an ideal world.

about “the probability of the null hypothesis” is as meaningless as “the colour of freedom”. It doesn’t have one!

For now, let’s suppose that you’re a Bayesian and that you are willing to interpret probability as a subjective thing: a degree of belief. Then we have no problems at all. I believe in some theories more strongly than others, so I’m perfectly entitled to assign probabilities to the event of “ H_0 is true”. So, as long as you think that probabilities are “in the mind” rather than “in the world”, you can base all of your statistical inference on Bayes’ theorem. Everything flows from this one simple equation. When we apply Bayes rule to the problem of hypothesis testing, the probability that the null hypothesis is true can be written like this:

$$P(H_0 \text{ is true} \mid \text{data } X) = \frac{P(\text{data } X \mid H_0 \text{ is true}) P(H_0 \text{ is true})}{P(\text{data } X)}$$

The probability $P(H_0 \text{ is true})$ is the *prior probability* of the null hypothesis: that is, the degree of confidence that you would have placed in the null hypothesis before observing your data. Then, once the data X are observed this prior probability is updated to a the *posterior probability* $P(H_0 \text{ is true} \mid \text{data } X)$, the degree of belief you assign to the null hypothesis now that you’ve looked at the data. The $P(\text{data } X \mid H_0 \text{ is true})$ term is called the *likelihood*, and it refers to the probability that you would have observed the data if the null hypothesis were true. The term $P(\text{data } X)$ describes the probability of the data set itself, and is actually pretty hard to calculate, though it’s not usually necessary to do so. Inference within the Bayesian framework involves a competition between different hypotheses. If we have two hypotheses to consider (H_0 and H_1), we would calculate *posterior odds ratio* that describes the relative plausibility of these two hypotheses:

$$\frac{P(H_1 \text{ is true} \mid \text{data } X)}{P(H_0 \text{ is true} \mid \text{data } X)}$$

If the posterior odds ratio is large, then we should prefer the alternative hypothesis to the null hypothesis; but if it is small, then we should prefer the null hypothesis. Obviously, I’m glossing over a lot of important details,¹⁵ but that’s the gist of the approach.

11.9.3 Traps

As you can see, the theory behind hypothesis testing is a mess, and even now there are arguments in statistics about how it “should” work. However, disagreements among statisticians are not our real concern here. Our real concern is practical data analysis. And while the “orthodox” approach to null hypothesis significance testing has many drawbacks, even an unrepentant Bayesian like myself would agree that they can be useful if used responsibly. Most of the time they give sensible answers, and you can use them to learn interesting things. Setting aside the various ideologies and historical confusions that we’ve discussed, the fact remains that the biggest danger in all of statistics is *thoughtlessness*. I don’t mean stupidity, here: I literally mean thoughtlessness. The rush to interpret a result without spending time thinking through what each test actually says about the data, and checking whether that’s consistent with how you’ve interpreted it. That’s where the biggest trap lies.

To give an example of this, consider the following example (see Gelman & Stern, 2006). Suppose I’m running my ESP study, and I’ve decided to analyse the data separately for the male participants and the female participants. Of the male participants, 33 out of 50 guessed the colour of the card correctly. This is a significant effect ($p = .03$). Of the female participants, 29 out of 50 guessed correctly. This is not a significant effect ($p = .32$). Upon observing this, it is extremely tempting for people to start wondering why there is a difference between males and females in terms of their psychic abilities. However, this is wrong. If you think about it, we haven’t *actually* run a test that explicitly compares males to females. All

¹⁵Not least of which is the fact that, if you specify silly hypotheses, Bayesian inference won’t be any more useful to you than orthodox analyses.

we have done is compare males to chance (binomial test was significant) and compared females to chance (binomial test was non significant). If we want to argue that there is a real difference between the males and the females, we should probably run a test of the null hypothesis that there is no difference! We can do that using a different hypothesis test,¹⁶ but when we do that it turns out that we have no evidence that males and females are significantly different ($p = .54$). Now do you think that there's anything fundamentally different between the two groups? Of course not. What's happened here is that the data from both groups (male and female) are pretty borderline: by pure chance, one of them happened to end up on the magic side of the $\alpha = .05$ line, and the other one didn't. That doesn't actually imply that males and females are different. This mistake is so common that you should always be wary of it: the difference between significant and not-significant is *not* evidence of a real difference – if you want to say that there's a difference between two groups, then you have to test for that difference!

The example above is just that: an example. I've singled it out because it's such a common one, but the bigger picture is that data analysis can be tricky to get right. Think about what it is you want to test, why you want to test it, and whether or not the answers that your test gives could possibly make any sense in the real world.

11.10 _____

Summary

- Research hypotheses and statistical hypotheses. Null and alternative hypotheses. (Section 11.1).
- Type 1 and Type 2 errors (Section 11.2)
- Test statistics and sampling distributions (Section 11.3)
- Hypothesis testing as a decision making process (Section 11.4)
- p -values as “soft” decisions (Section 11.5)
- Writing up the results of a hypothesis test (Section 11.6)
- Effect size and power (Section 11.8)
- A few issues to consider regarding hypothesis testing (Section 11.9)

¹⁶In this case, the Pearson chi-square test of independence (Chapter 12; `chisq.test()` in R) is what we use; see also the `prop.test()` function.

Part V.

Statistical tools

12. Categorical data analysis

Now that we've got the basic theory behind hypothesis testing, it's time to start looking at specific tests that are commonly used in psychology. So where should we start? Not every textbook agrees on where to start, but I'm going to start with " χ^2 tests" (this chapter) and "*t*-tests" (Chapter 13). Both of these tools are very frequently used in scientific practice, and while they're not as powerful as "analysis of variance" (Chapter 14) and "regression" (Chapter 15) they're much easier to understand.

The term "categorical data" is just another name for "nominal scale data". It's nothing that we haven't already discussed, it's just that in the context of data analysis people tend to use the term "categorical data" rather than "nominal scale data". I don't know why. In any case, **categorical data analysis** refers to a collection of tools that you can use when your data are nominal scale. However, there are a lot of different tools that can be used for categorical data analysis, and this chapter only covers a few of the more common ones.

12.1

The χ^2 goodness-of-fit test

The Pearson χ^2 goodness-of-fit test is one of the oldest hypothesis tests around: it was invented by Karl Pearson around the turn of the century (Pearson, 1900), with some corrections made later by Sir Ronald Fisher (Fisher, 1922). To introduce the statistical problem that it addresses, let's start with some psychology...

12.1.1 The cards data

Over the years, there have been a lot of studies showing that humans have a lot of difficulties in simulating randomness. Try as we might to "act" random, we *think* in terms of patterns and structure, and so when asked to "do something at random", what people actually do is anything but random. As a consequence, the study of human randomness (or non-randomness, as the case may be) opens up a lot of deep psychological questions about how we think about the world. With this in mind, let's consider a very simple study. Suppose I asked people to imagine a shuffled deck of cards, and mentally pick one card from this imaginary deck "at random". After asking, say, $N = 200$ people to do this, I'd like to look at the data and figure out whether or not the cards that people pretended to select were really random.¹ How can I do this? In order to answer this question, we'll need to have a look at my data, which I've saved in an R data file called `cards.Rdata`. Let's take a look:

¹I've run experiments like this, and this example is loosely based on the real data.

```

> load( "cards.Rdata" )
> who()
-- Name --    -- Class --    -- Size --
suit.choice   factor        200

```

There's a single variable in the data file called `suitchoice`, and it's a factor. If we use the `head()` function to get R to show us the first few entries, and the `summary()` function to get an overview of what the data look like, here's what we get:

```

> # show the first six observations:
> head( suit.choice )
[1] spades diamonds hearts spades hearts clubs
Levels: clubs diamonds hearts spades

> # print out a summary of the observations:
> summary( suit.choice )
clubs diamonds hearts spades
 35      51      64      50

```

That little frequency table is quite helpful. Looking at it, there's a bit of a hint that people might be more likely to select hearts than clubs, but it's not completely obvious whether that's true or not. So we'll probably have to do some kind of statistical analysis to find out, which is what I'm going to talk about in the next section. However, because the statistical methods that we're going to look at are based on the analysis of *frequency tables* like the one that the `summary()` function printed out, we really ought to construct a table of **observed frequencies** and store it as a variable. Let's do that now, using the `table()` function:²

```

> O <- table( suit.choice )  # construct the table
> print( O )    # print it out
suit.choice
clubs diamonds hearts spades
 35      51      64      50

```

That being said, it's worth keeping in mind that the `table()` function actually creates a slightly complicated object. That is, there's a few things hidden inside this `O` variable that might confuse you if you start playing around with it. So, what I'm going to do for our example is this:

```

> O <- tabulate( suit.choice )
> names(O) <- c("clubs","diamonds","hearts","spades")
> print(O)
clubs diamonds hearts spades
 35      51      64      50

```

The output looks almost the same, but there's nothing fancy going on behind the scenes. Our variable `O` is just an ordinary numeric vector, of the kind that we've been using throughout the book so far.

Excellent. From this point on, we'll treat this table as the data that we're looking to analyse. However, since I'm going to have to talk about this data in mathematical terms (sorry!) it might be a good idea to be clear about what the notation is. In R, if I wanted to pull out the number of people that selected diamonds, I could do it by name by typing `O["diamonds"]` but, since "`diamonds`" is second element of the

²Normally you'd try to use a more meaningful name to describe the variable, such as `suit.counts` or something. However, since the mathematical notation used for the observed frequency is O , I thought it might be useful to name the R variable `O`, so that the R commands that you see me typing look as similar as possible to the mathematical formulas that they calculate.

`o` vector, it's equally effective to refer to it as `o[2]`. The mathematical notation that we use is pretty much identical to this convention: but instead of writing `o[2]`, we write it O_2 . The relationship between the English descriptions, the R commands, and the mathematical symbols are illustrated below:

label	index, i	math. symbol	R command	the value
clubs, ♣	1	O_1	<code>o[1]</code>	35
diamonds, ♦	2	O_2	<code>o[2]</code>	51
hearts, ♥	3	O_3	<code>o[3]</code>	64
spades, ♠	4	O_4	<code>o[4]</code>	50

Hopefully that's pretty clear: in general we refer to the number of observations that fall in the i -th category by the symbol O_i . The only other thing that I'd point out is that statisticians often like to group together all of the O_i values into a vector, which I'll refer to as \mathbf{O} .

$$\mathbf{O} = (O_1, O_2, O_3, O_4)$$

Again, there's nothing new or interesting here: it's just notation. If I say that $\mathbf{O} = (35, 51, 64, 50)$ all I'm doing is describing the table of observed frequencies `o`, but I'm referring to it using mathematical notation, rather than by referring to an R variable.

12.1.2 The null hypothesis and the alternative hypothesis

As the last section indicated, our research hypothesis is that “people don’t choose cards randomly”. What we’re going to want to do now is translate this into some statistical hypotheses, and construct a statistical test of those hypotheses. The test that I’m going to describe to you is **Pearson’s χ^2 goodness of fit test**, and as is so often the case, we have to begin by carefully constructing our null hypothesis. In this case, it’s pretty easy. First, let’s state the null hypothesis in words:

$$H_0: \quad \text{All four suits are chosen with equal probability}$$

Now, because this is statistics, we have to be able to say the same thing in a mathematical way. To do this, let’s use the notation P_j to refer to the true probability that the j -th suit is chosen. If the null hypothesis is true, then each of the four suits has a 25% chance of being selected: in other words, our null hypothesis claims that $P_1 = .25$, $P_2 = .25$, $P_3 = .25$ and finally that $P_4 = .25$. However, in the same way that we can group our observed frequencies into a vector \mathbf{O} that summarises the entire data set, we can use \mathbf{P} to refer to the probabilities that correspond to our null hypothesis. So if I let the vector $\mathbf{P} = (P_1, P_2, P_3, P_4)$ refer to the collection of probabilities that describe our null hypothesis, then we have

$$H_0: \quad \mathbf{P} = (.25, .25, .25, .25)$$

In this particular instance, our null hypothesis corresponds to a vector of probabilities \mathbf{P} in which all of the probabilities are equal to one another. But this doesn’t have to be the case. For instance, if the experimental task was for people to imagine they were drawing from a deck that had twice as many clubs as any other suit, then the null hypothesis would correspond to something like $\mathbf{P} = (.4, .2, .2, .2)$. As long as the probabilities are all positive numbers, and they all sum to 1, then it’s a perfectly legitimate choice for the null hypothesis. However, the most common use of the goodness of fit test is to test a null hypothesis that all of the categories are equally likely, so we’ll stick to that for our example.

What about our alternative hypothesis, H_1 ? All we’re really interested in is demonstrating that the probabilities involved aren’t all identical (that is, people’s choices weren’t completely random). As a consequence, the “human friendly” versions of our hypotheses look like this:

$$H_0: \quad \text{All four suits are chosen with equal probability}$$

$$H_1: \quad \text{At least one of the suit-choice probabilities } \textit{isn't} .25$$

and the “mathematician friendly” version is

$$\begin{aligned} H_0: \quad & \mathbf{P} = (.25, .25, .25, .25) \\ H_1: \quad & \mathbf{P} \neq (.25, .25, .25, .25) \end{aligned}$$

Conveniently, the mathematical version of the hypotheses looks quite similar to an R command defining a vector. So maybe what I should do is store the \mathbf{P} vector in R as well, since we’re almost certainly going to need it later. And because I’m Mister Imaginative, I’ll call this R vector \mathbf{P} ,

```
> P <- c(.25, .25, .25, .25) # enter the probabilities
> names(P) <- names(O)      # copy the labels across from O to P
> print(P)
  clubs diamonds   hearts   spades
  0.25      0.25      0.25      0.25
```

12.1.3 The “goodness of fit” test statistic

At this point, we have our observed frequencies \mathbf{O} and a collection of probabilities \mathbf{P} that describes the null hypothesis that we want to test. We’ve stored these in R as the corresponding variables \mathbf{O} and \mathbf{P} . What we now want to do is construct a test of the null hypothesis. As always, if we want to test H_0 against H_1 , we’re going to need a test statistic. The basic trick that a “goodness of fit” test uses is to construct a test statistic that measures how “close” the data are to the null hypothesis. If the data don’t resemble what you’d “expect” to see if the null hypothesis were true, then it probably isn’t true. Okay, if the null hypothesis were true, what would we expect to see? Or, to use the correct terminology, what are the **expected frequencies**. There are $N = 200$ observations, and (if the null is true) the probability of any one of them choosing a heart is $P_3 = .25$, so I guess we’re expecting $200 \times .25 = 50$ hearts, right? Or, more specifically, if we let E_i refer to “the number of category i responses that we’re expecting if the null is true”, then

$$E_i = N \times P_i$$

This is pretty easy to calculate in R:

```
> N <- 200      # store the sample size
> E <- N * P    # calculate the expected frequencies
> print(E)      # and now, let's have a look...
  clubs diamonds   hearts   spades
  50      50      50      50
```

None of which is very surprising.

Now, how do we translate this into a test statistic? Clearly, what we want to do is compare the *expected* number of observations in each category (E_i) with the *observed* number of observations in that category (O_i). And on the basis of this comparison, we ought to be able to come up with a good test statistic. To start with, let’s calculate the difference between what the null hypothesis expected us to find and what we actually did find. That is, we calculate the “observed minus expected” difference score, $O_i - E_i$. This is illustrated in the following table.

		♣	♦	♥	♠
expected frequency	E_i	50	50	50	50
observed frequency	O_i	35	51	64	50
difference score	$O_i - E_i$	-15	1	14	0

The same calculations can be done in R, using our \mathbf{E} and \mathbf{O} variables:

```
> O-E
clubs diamonds hearts spades
-15      1      14      0
```

Regardless of whether we do the calculations by hand or whether we do them in R, it's clear that people chose more hearts and fewer clubs than the null hypothesis predicted. However, a moment's thought suggests that these raw differences aren't quite what we're looking for. Intuitively, it feels like it's just as bad when the null hypothesis predicts too few observations (which is what happened with hearts) as it is when it predicts too many (which is what happened with clubs). So it's a bit weird that we have a negative number for hearts and a positive number for clubs. One easy way to fix this is to square everything, so that we now calculate the squared differences, $(E_i - O_i)^2$. As before, we could do this by hand, but it's easier to do it in R...

```
> (O-E)^2
clubs diamonds hearts spades
225      1     196      0
```

Now we're making progress. What we've got now is a collection of numbers that are big whenever the null hypothesis makes a bad prediction (clubs and hearts), but are small whenever it makes a good one (diamonds and spades). Next, for some technical reasons that I'll explain in a moment, let's also divide all these numbers by the expected frequency E_i , so we're actually calculating $\frac{(E_i - O_i)^2}{E_i}$. Since $E_i = 50$ for all categories in our example, it's not a very interesting calculation, but let's do it anyway. The R command becomes:

```
> (O-E)^2 / E
clubs diamonds hearts spades
4.50      0.02    3.92      0.00
```

In effect, what we've got here are four different “error” scores, each one telling us how big a “mistake” the null hypothesis made when we tried to use it to predict our observed frequencies. So, in order to convert this into a useful test statistic, one thing we could do is just add these numbers up. The result is called the **goodness of fit** statistic, conventionally referred to either as X^2 or GOF. We can calculate it using this command in R

```
> sum( (O-E)^2 / E )
[1] 8.44
```

The formula for this statistic looks remarkably similar to the R command. If we let k refer to the total number of categories (i.e., $k = 4$ for our cards data), then the X^2 statistic is given by:

$$X^2 = \sum_{i=1}^k \frac{(O_i - E_i)^2}{E_i}$$

Intuitively, it's clear that if X^2 is small, then the observed data O_i are very close to what the null hypothesis predicted E_i , so we're going to need a large X^2 statistic in order to reject the null. As we've seen from our calculations, in our cards data set we've got a value of $X^2 = 8.44$. So now the question becomes, is this a big enough value to reject the null?

12.1.4 The sampling distribution of the GOF statistic

To determine whether or not a particular value of X^2 is large enough to justify rejecting the null hypothesis, we're going to need to figure out what the sampling distribution for X^2 would be if the null

hypothesis were true. So that's what I'm going to do in this section. I'll show you in a fair amount of detail how this sampling distribution is constructed, and then – in the next section – use it to build up a hypothesis test. If you want to cut to the chase and are willing to take it on faith that the sampling distribution is a **chi-squared (χ^2) distribution** with $k - 1$ degrees of freedom, you can skip the rest of this section. However, if you want to understand *why* the goodness of fit test works the way it does, read on...

Okay, let's suppose that the null hypothesis is actually true. If so, then the true probability that an observation falls in the i -th category is P_i – after all, that's pretty much the definition of our null hypothesis. Let's think about what this actually means. If you think about it, this is kind of like saying that "nature" makes the decision about whether or not the observation ends up in category i by flipping a weighted coin (i.e., one where the probability of getting a head is P_j). And therefore, we can think of our observed frequency O_i by imagining that nature flipped N of these coins (one for each observation in the data set)... and exactly O_i of them came up heads. Obviously, this is a pretty weird way to think about the experiment. But what it does (I hope) is remind you that we've actually seen this scenario before. It's exactly the same set up that gave rise to the binomial distribution in Section 9.3. In other words, if the null hypothesis is true, then it follows that our observed frequencies were generated by sampling from a binomial distribution:

$$O_i \sim \text{Binomial}(P_i, N)$$

Now, if you remember from our discussion of the central limit theorem (Section 10.4), the binomial distribution starts to look pretty much identical to the normal distribution, especially when N is large and when P_i isn't *too* close to 0 or 1. In other words as long as $N \times P_i$ is large enough – or, to put it another way, when the expected frequency E_i is large enough – the theoretical distribution of O_i is approximately normal. Better yet, if O_i is normally distributed, then so is $(O_i - E_i)/\sqrt{E_i}$... since E_i is a fixed value, subtracting off E_i and dividing by $\sqrt{E_i}$ changes the mean and standard deviation of the normal distribution; but that's all it does. Okay, so now let's have a look at what our goodness of fit statistic actually *is*. What we're doing is taking a bunch of things that are normally-distributed, squaring them, and adding them up. Wait. We've seen that before too! As we discussed in Section 9.5, when you take a bunch of things that have a standard normal distribution (i.e., mean 0 and standard deviation 1), square them, then add them up, then the resulting quantity has a chi-square distribution. So now we know that the null hypothesis predicts that the sampling distribution of the goodness of fit statistic is a chi-square distribution. Cool.

There's one last detail to talk about, namely the degrees of freedom. If you remember back to Section 9.5, I said that if the number of things you're adding up is k , then the degrees of freedom for the resulting chi-square distribution is k . Yet, what I said at the start of this section is that the actual degrees of freedom for the chi-square goodness of fit test is $k - 1$. What's up with that? The answer here is that what we're supposed to be looking at is the number of genuinely *independent* things that are getting added together. And, as I'll go on to talk about in the next section, even though there's k things that we're adding, only $k - 1$ of them are truly independent; and so the degrees of freedom is actually only $k - 1$. That's the topic of the next section.³

12.1.5 Degrees of freedom

When I introduced the chi-square distribution in Section 9.5, I was a bit vague about what "**degrees of freedom**" actually *means*. Obviously, it matters: looking Figure 12.1 you can see that if we change the

³I should point out that this issue does complicate the story somewhat: I'm not going to cover it in this book, but there's a sneaky trick that you can do to rewrite the equation for the goodness of fit statistic as a sum over $k - 1$ independent things. When we do so we get the "proper" sampling distribution, which is chi-square with $k - 1$ degrees of freedom. In fact, in order to get the maths to work out properly, you actually have to rewrite things that way. But it's beyond the scope of an introductory book to show the maths in that much detail: all I wanted to do is give you a sense of why the goodness of fit statistic is associated with the chi-squared distribution.

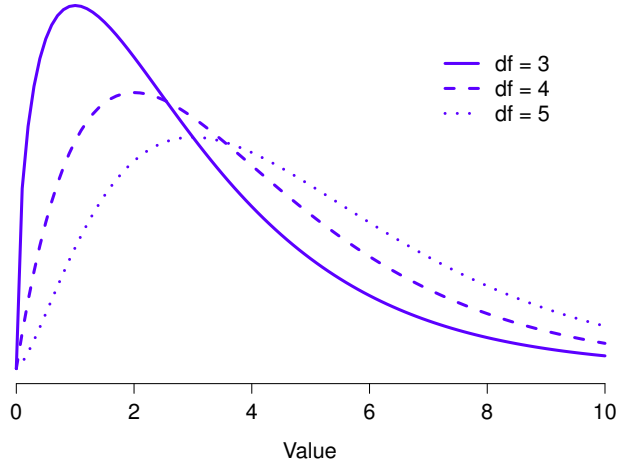


Figure 12.1: Chi-square distributions with different values for the “degrees of freedom”.

degrees of freedom, then the chi-square distribution changes shape quite substantially. But what exactly *is* it? Again, when I introduced the distribution and explained its relationship to the normal distribution, I did offer an answer... it’s the number of “normally distributed variables” that I’m squaring and adding together. But, for most people, that’s kind of abstract, and not entirely helpful. What we really need to do is try to understand degrees of freedom in terms of our data. So here goes.

The basic idea behind degrees of freedom is quite simple: you calculate it by counting up the number of distinct “quantities” that are used to describe your data; and then subtracting off all of the “constraints” that those data must satisfy.⁴ This is a bit vague, so let’s use our cards data as a concrete example. We describe our data using four numbers, O_1 , O_2 , O_3 and O_4 corresponding to the observed frequencies of the four different categories (hearts, clubs, diamonds, spades). These four numbers are the *random outcomes* of our experiment. But, my experiment actually has a fixed constraint built into it: the sample size N .⁵ That is, if we know how many people chose hearts, how many chose diamonds and how many chose clubs; then we’d be able to figure out exactly how many chose spades. In other words, although our data are described using four numbers, they only actually correspond to $4 - 1 = 3$ degrees of freedom. A slightly different way of thinking about it is to notice that there are four *probabilities* that we’re interested in (again, corresponding to the four different categories), but these probabilities must sum to one, which imposes a constraint. Therefore, the degrees of freedom is $4 - 1 = 3$. Regardless of whether you want to think about it in terms of the observed frequencies or in terms of the probabilities, the answer is the same. In general, when running the chi-square goodness of fit test for an experiment involving k groups,

⁴I feel obliged to point out that this is an over-simplification. It works nicely for quite a few situations; but every now and then we’ll come across degrees of freedom values that aren’t whole numbers. Don’t let this worry you too much – when you come across this, just remind yourself that “degrees of freedom” is actually a bit of a messy concept, and that the nice simple story that I’m telling you here isn’t the whole story. For an introductory class, it’s usually best to stick to the simple story: but I figure it’s best to warn you to expect this simple story to fall apart. If I didn’t give you this warning, you might start getting confused when you see $df = 3.4$ or something; and (incorrectly) thinking that you had misunderstood something that I’ve taught you, rather than (correctly) realising that there’s something that I haven’t told you.

⁵In practice, the sample size isn’t always fixed... e.g., we might run the experiment over a fixed period of time, and the number of people participating depends on how many people show up. That doesn’t matter for the current purposes.

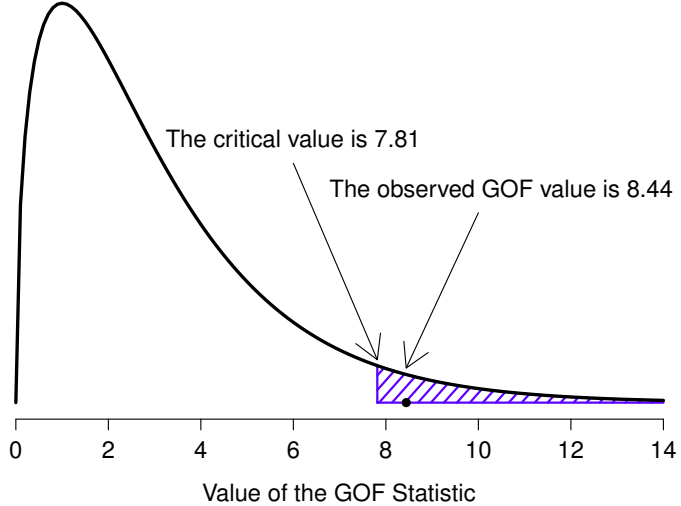


Figure 12.2: Illustration of how the hypothesis testing works for the chi-square goodness of fit test.

then the degrees of freedom will be $k - 1$.

12.1.6 Testing the null hypothesis

The final step in the process of constructing our hypothesis test is to figure out what the rejection region is. That is, what values of X^2 would lead us to reject the null hypothesis. As we saw earlier, large values of X^2 imply that the null hypothesis has done a poor job of predicting the data from our experiment, whereas small values of X^2 imply that it's actually done pretty well. Therefore, a pretty sensible strategy would be to say there is some critical value, such that if X^2 is bigger than the critical value we reject the null; but if X^2 is smaller than this value we retain the null. In other words, to use the language we introduced in Chapter 11 the chi-squared goodness of fit test is always a **one-sided test**. Right, so all we have to do is figure out what this critical value is. And it's pretty straightforward. If we want our test to have significance level of $\alpha = .05$ (that is, we are willing to tolerate a Type I error rate of 5%), then we have to choose our critical value so that there is only a 5% chance that X^2 could get to be that big if the null hypothesis is true. That is to say, we want the 95th percentile of the sampling distribution. This is illustrated in Figure 12.2.

Ah, but – I hear you ask – how do I calculate the 95th percentile of a chi-squared distribution with $k - 1$ degrees of freedom? If only R had some function, called... oh, I don't know, `qchisq()` ... that would let you calculate this percentile (see Chapter 9 if you've forgotten). Like this...

```
> qchisq( p = .95, df = 3 )
[1] 7.814728
```

So if our X^2 statistic is bigger than 7.81 or so, then we can reject the null hypothesis. Since we actually calculated that before (i.e., $X^2 = 8.44$) we can reject the null. If we want an exact p -value, we can calculate it using the `pchisq()` function:

```
> pchisq( q = 8.44, df = 3, lower.tail = FALSE )
[1] 0.03774185
```

This is hopefully pretty straightforward, as long as you recall that the “`p`” form of the probability distribution functions in R always calculates the probability of getting a value of *less* than the value you entered (in this case 8.44). We want the opposite: the probability of getting a value of 8.44 or *more*. That’s why I told R to use the upper tail, not the lower tail. That said, it’s usually easier to calculate the *p*-value this way:

```
> 1-pchisq( q = 8.44, df = 3 )
[1] 0.03774185
```

So, in this case we would reject the null hypothesis, since $p < .05$. And that’s it, basically. You now know “Pearson’s χ^2 test for the goodness of fit”. Lucky you.

12.1.7 Doing the test in R

Gosh darn it. Although we did manage to do everything in R as we were going through that little example, it does rather feel as if we’re typing too many things into the magic computing box. And I *hate* typing. Fortunately, R provides a function called `chisq.test()` which will do all the calculations for you. If you’re trying to do a chi-square goodness of fit test, then there’s two arguments you need to specify: the data, `x`, which needs to be a vector containing the observed probabilities (i.e., `o`), and the probabilities `p` that specify the null hypothesis (in our case, `P`). So the command you’re looking for is something like this:

```
> chisq.test( x = o, p = P )
```

However, if you don’t bother to specify `p`, R will assume that the null hypothesis is that all of the categories are equally likely. Since that actually is our null hypothesis in this case, there’s no need to bother specifying the `p` argument. So let’s have a look at the `chisq.test()` function, the way you’d actually use it in real life to do a chi-square goodness of fit test. First, in real life you’d use a better name than `o` to describe your data, so let’s actually do that:

```
> suit.table <- table( suit.choice )
```

And then you would run your hypothesis test like this:

```
> chisq.test( suit.table )

Chi-squared test for given probabilities

data: suit.table
X-squared = 8.44, df = 3, p-value = 0.03774
```

As you can see, these are the exact same numbers we calculated previously. So you’re done. In short, to run a chi-square goodness of fit test, what you do is construct the table of observed frequencies (using one of the tabulation functions discussed in Section 7.1), and then use `chisq.test()` to run the chi-square test itself.

12.1.8 How to report the results of the test

So now you know how the test works, and you know how to do the test using a wonderful magic computing box. The next thing you need to know is how to write up the results. After all, there's no point in designing and running an experiment and then analysing the data if you don't tell anyone about it! So let's now talk about what you need to do when reporting your analysis. Let's stick with our card-suits example. If I wanted to write this result up for a paper or something, the conventional way to report this would be to write something like this:

Of the 200 participants in the experiment, 64 selected hearts, 51 selected diamonds, 50 selected spades, and 35 selected clubs. A chi-square goodness of fit was conducted to test whether the choice probabilities were identical for all four suits. The results were significant ($\chi^2(3) = 8.44, p < .05$), suggesting that people did not select suits purely at random.

This is pretty straightforward, and hopefully it seems pretty unremarkable. That said, there's a few things that you should note about this description:

- *The statistical test is preceded by the descriptive statistics.* That is, I told the reader something about what the data look like before going on to do the test. In general, this is good practice: always remember that your reader doesn't know your data anywhere near as well as you do. So unless you describe it to them properly, the statistical tests won't make any sense to them, and they'll get frustrated and cry.
- *The description tells you what the null hypothesis being tested is.* To be honest, writers don't always do this, but it's often a good idea in those situations where some ambiguity exists; or when you can't rely on your readership being intimately familiar with the statistical tools that you're using. Quite often the reader might not know (or remember) all the details of the test that you're using, so it's a kind of politeness to "remind" them! As far as the goodness of fit test goes, you can usually rely on a scientific audience knowing how it works (since it's covered in most intro stats classes). However, it's still a good idea to be explicit about stating the null hypothesis (briefly!) because the null hypothesis can be different depending on what you're using the test for. For instance, in the cards example my null hypothesis was that all the four suit probabilities were identical (i.e., $P_1 = P_2 = P_3 = P_4 = 0.25$), but there's nothing special about that hypothesis. I could just as easily have tested the null hypothesis that $P_1 = 0.7$ and $P_2 = P_3 = P_4 = 0.1$ using a goodness of fit test. So it's helpful to the reader if you explain to them what your null hypothesis was. Also, notice that I described the null hypothesis in words, not in maths. That's perfectly acceptable. You can describe it in maths if you like, but since most readers find words easier to read than symbols, most writers tend to describe the null using words if they can.
- *A "stat block" is included.* When reporting the results of the test itself, I didn't just say that the result was significant, I included a "stat block" (i.e., the dense mathematical-looking part in the parentheses), which reports all the "raw" statistical data. For the chi-square goodness of fit test, the information that gets reported is the test statistic (that the goodness of fit statistic was 8.44), the information about the distribution used in the test (χ^2 with 3 degrees of freedom, which is usually shortened to $\chi^2(3)$), and then the information about whether the result was significant (in this case $p < .05$). The particular information that needs to go into the stat block is different for every test, and so each time I introduce a new test I'll show you what the stat block should look like.⁶ However the general principle is that you should always provide enough information so that the reader could check the test results themselves if they really wanted to.

⁶Well, sort of. The conventions for how statistics should be reported tend to differ somewhat from discipline to discipline; I've tended to stick with how things are done in psychology, since that's what I do. But the general principle of providing enough information to the reader to allow them to check your results is pretty universal, I think.

- *The results are interpreted.* In addition to indicating that the result was significant, I provided an interpretation of the result (i.e., that people didn't choose randomly). This is also a kindness to the reader, because it tells them something about what they should believe about what's going on in your data. If you don't include something like this, it's really hard for your reader to understand what's going on.⁷

As with everything else, your overriding concern should be that you *explain* things to your reader. Always remember that the point of reporting your results is to communicate to another human being. I cannot tell you just how many times I've seen the results section of a report or a thesis or even a scientific article that is just gibberish, because the writer has focused solely on making sure they've included all the numbers, and forgotten to actually communicate with the human reader.

12.1.9 A comment on statistical notation

Satan delights equally in statistics and in quoting scripture

– H.G. Wells

If you've been reading very closely, and are as much of a mathematical pedant as I am, there is one thing about the way I wrote up the chi-square test in the last section that might be bugging you a little bit. There's something that feels a bit wrong with writing " $\chi^2(3) = 8.44$ ", you might be thinking. After all, it's the goodness of fit statistic that is equal to 8.44, so shouldn't I have written $X^2 = 8.44$ or maybe $GOF = 8.44$? This seems to be conflating the *sampling distribution* (i.e., χ^2 with $df = 3$) with the *test statistic* (i.e., X^2). Odds are you figured it was a typo, since χ and X look pretty similar. Oddly, it's not. Writing $\chi^2(3) = 8.44$ is essentially a highly condensed way of writing "the sampling distribution of the test statistic is $\chi^2(3)$, and the value of the test statistic is 8.44".

In one sense, this is kind of stupid. There are *lots* of different test statistics out there that turn out to have a chi-square sampling distribution: the X^2 statistic that we've used for our goodness of fit test is only one of many (albeit one of the most commonly encountered ones). In a sensible, perfectly organised world, we'd *always* have a separate name for the test statistic and the sampling distribution: that way, the stat block itself would tell you exactly what it was that the researcher had calculated. Sometimes this happens. For instance, the test statistic used in the Pearson goodness of fit test is written X^2 ; but there's a closely related test known as the *G-test*⁸ (Sokal & Rohlf, 1994), in which the test statistic is written as G . As it happens, the Pearson goodness of fit test and the *G-test* both test the same null hypothesis; and the sampling distribution is exactly the same (i.e., chi-square with $k - 1$ degrees of freedom). If I'd done a *G-test* for the cards data rather than a goodness of fit test, then I'd have ended up with a test statistic of $G = 8.65$, which is slightly different from the $X^2 = 8.44$ value that I got earlier; and produces a slightly smaller p -value of $p = .034$. Suppose that the convention was to report the test statistic, then the sampling distribution, and then the p -value. If that were true, then these two situations would produce different stat blocks: my original result would be written $X^2 = 8.44, \chi^2(3), p = .038$, whereas the new version using the *G-test* would be written as $G = 8.65, \chi^2(3), p = .034$. However, using the

⁷To some people, this advice might sound odd, or at least in conflict with the “usual” advice on how to write a technical report. Very typically, students are told that the “results” section of a report is for describing the data and reporting statistical analysis; and the “discussion” section is for providing interpretation. That’s true as far as it goes, but I think people often interpret it way too literally. The way I usually approach it is to provide a quick and simple interpretation of the data in the results section, so that my reader understands what the data are telling us. Then, in the discussion, I try to tell a bigger story; about how my results fit with the rest of the scientific literature. In short; don’t let the “interpretation goes in the discussion” advice turn your results section into incomprehensible garbage. Being understood by your reader is *much* more important.

⁸Complicating matters, the *G-test* is a special case of a whole class of tests that are known as *likelihood ratio tests*. I don’t cover LRTs in this book, but they are quite handy things to know about.

condensed reporting standard, the original result is written $\chi^2(3) = 8.44, p = .038$, and the new one is written $\chi^2(3) = 8.65, p = .034$, and so it's actually unclear which test I actually ran.

So why don't we live in a world in which the contents of the stat block uniquely specifies what tests were ran? The deep reason is that life is messy. We (as users of statistical tools) want it to be nice and neat and organised... we want it to be *designed*, as if it were a product. But that's not how life works: statistics is an intellectual discipline just as much as any other one, and as such it's a massively distributed, partly-collaborative and partly-competitive project that no-one really understands completely. The things that you and I use as data analysis tools weren't created by an Act of the Gods of Statistics; they were invented by lots of different people, published as papers in academic journals, implemented, corrected and modified by lots of other people, and then explained to students in textbooks by someone else. As a consequence, there's a *lot* of test statistics that don't even have names; and as a consequence they're just given the same name as the corresponding sampling distribution. As we'll see later, any test statistic that follows a χ^2 distribution is commonly called a "chi-square statistic"; anything that follows a t -distribution is called a " t -statistic" and so on. But, as the X^2 versus G example illustrates, two different things with the same sampling distribution are still, well, different.

As a consequence, it's sometimes a good idea to be clear about what the actual test was that you ran, especially if you're doing something unusual. If you just say "chi-square test", it's not actually clear what test you're talking about. Although, since the two most common chi-square tests are the goodness of fit test and the independence test (Section 12.2), most readers with stats training can probably guess. Nevertheless, it's something to be aware of.

12.2

The χ^2 test of independence

GUARDBOT 1: *Halt!*
GUARDBOT 2: *Be you robot or human?*
LEELA: *Robot...we be.*
FRY: *Uh, yup! Just two robots out roboting it up! Eh?*
GUARDBOT 1: *Administer the test.*
GUARDBOT 2: *Which of the following would you most prefer?*
 A: A puppy, B: A pretty flower from your sweetie,
 or C: A large properly-formatted data file?
GUARDBOT 1: *Choose!*
 – Futurama, “Fear of a Bot Planet”

The other day I was watching an animated documentary examining the quaint customs of the natives of the planet *Chapek 9*. Apparently, in order to gain access to their capital city, a visitor must prove that they're a robot, not a human. In order to determine whether or not visitor is human, they ask whether the visitor prefers puppies, flowers or large, properly formatted data files. "Pretty clever," I thought to myself "but what if humans and robots have the same preferences? That probably wouldn't be a very good test then, would it?" As it happens, I got my hands on the testing data that the civil authorities of *Chapek 9* used to check this. It turns out that what they did was very simple... they found a bunch of robots and a bunch of humans and asked them what they preferred. I saved their data in a file called *chapek9.Rdata*, which I can now load and have a quick look at:

```
> load("chapek9.Rdata")
> who(TRUE)
-- Name -- -- Class -- -- Size --
```

```
chapek9      data.frame   180 x 2
$species     factor       180
$choice      factor       180
```

Okay, so we have a single data frame called `chapek9`, which contains two factors, `species` and `choice`. As always, it's nice to have a quick look at the data,

```
> head(chapek9)
  species choice
1  robot   flower
2  human   data
3  human   data
4  human   data
5  robot   data
6  human   flower
```

and then take a `summary()`,

```
> summary(chapek9)
  species     choice
robot:87    puppy : 28
human:93    flower: 43
            data  :109
```

In total there are 180 entries in the data frame, one for each person (counting both robots and humans as “people”) who was asked to make a choice. Specifically, there's 93 humans and 87 robots; and overwhelmingly the preferred choice is the data file. However, these summaries don't address the question we're interested in. To do that, we need a more detailed description of the data. What we want to do is look at the `choices` broken down by `species`. That is, we need to cross-tabulate the data (see Section 7.1). There's quite a few ways to do this, as we've seen, but since our data are stored in a data frame, it's convenient to use the `xtabs()` function.

```
> chapek9.xtabs <- xtabs(~ choice + species, data = chapek9)
> print(chapek9.xtabs )
  species
choice  robot human
puppy    13    15
flower   30    13
data     44    65
```

That's more or less what we're after. So, if we add the row and column totals (which is convenient for the purposes of explaining the statistical tests), we would have a table like this,

	Robot	Human	Total
Puppy	13	15	28
Flower	30	13	43
Data file	44	65	109
Total	87	93	180

which actually would be a nice way to report the descriptive statistics for this data set. In any case, it's quite clear that the vast majority of the humans chose the data file, whereas the robots tended to be a lot more even in their preferences. Leaving aside the question of *why* the humans might be more likely to choose the data file for the moment (which does seem quite odd, admittedly), our first order of business is to determine if the discrepancy between human choices and robot choices in the data set is statistically significant.

12.2.1 Constructing our hypothesis test

How do we analyse this data? Specifically, since my *research* hypothesis is that “humans and robots answer the question in different ways”, how can I construct a test of the *null* hypothesis that “humans and robots answer the question the same way”? As before, we begin by establishing some notation to describe the data:

	Robot	Human	Total
Puppy	O_{11}	O_{12}	R_1
Flower	O_{21}	O_{22}	R_2
Data file	O_{31}	O_{32}	R_3
Total	C_1	C_2	N

In this notation we say that O_{ij} is a count (observed frequency) of the number of respondents that are of species j (robots or human) who gave answer i (puppy, flower or data) when asked to make a choice. The total number of observations is written N , as usual. Finally, I’ve used R_i to denote the row totals (e.g., R_1 is the total number of people who chose the flower), and C_j to denote the column totals (e.g., C_1 is the total number of robots).⁹

So now let’s think about what the null hypothesis says. If robots and humans are responding in the same way to the question, it means that the probability that “a robot says puppy” is the same as the probability that “a human says puppy”, and so on for the other two possibilities. So, if we use P_{ij} to denote “the probability that a member of species j gives response i ” then our null hypothesis is that:

- H_0 : All of the following are true:
 $P_{11} = P_{12}$ (same probability of saying “puppy”),
 $P_{21} = P_{22}$ (same probability of saying “flower”), and
 $P_{31} = P_{32}$ (same probability of saying “data”).

And actually, since the null hypothesis is claiming that the true choice probabilities don’t depend on the species of the person making the choice, we can let P_i refer to this probability: e.g., P_1 is the true probability of choosing the puppy.

Next, in much the same way that we did with the goodness of fit test, what we need to do is calculate the expected frequencies. That is, for each of the observed counts O_{ij} , we need to figure out what the null hypothesis would tell us to expect. Let’s denote this expected frequency by E_{ij} . This time, it’s a little bit trickier. If there are a total of C_j people that belong to species j , and the true probability of anyone (regardless of species) choosing option i is P_i , then the expected frequency is just:

$$E_{ij} = C_j \times P_i$$

Now, this is all very well and good, but we have a problem. Unlike the situation we had with the goodness of fit test, the null hypothesis doesn’t actually specify a particular value for P_i . It’s something we have to estimate (Chapter 10) from the data! Fortunately, this is pretty easy to do. If 28 out of 180 people selected the flowers, then a natural estimate for the probability of choosing flowers is 28/180, which is approximately .16. If we phrase this in mathematical terms, what we’re saying is that our estimate for the probability of choosing option i is just the row total divided by the total sample size:

$$\hat{P}_i = \frac{R_i}{N}$$

⁹A technical note. The way I’ve described the test pretends that the column totals are fixed (i.e., the researcher intended to survey 87 robots and 93 humans) and the row totals are random (i.e., it just turned out that 28 people chose the puppy). To use the terminology from my mathematical statistics textbook (Hogg, McKean, & Craig, 2005) I should technically refer to this situation as a chi-square test of homogeneity; and reserve the term chi-square test of independence for the situation where both the row and column totals are random outcomes of the experiment. In the initial drafts of this book that’s exactly what I did. However, it turns out that these two tests are identical; and so I’ve collapsed them together.

Therefore, our expected frequency can be written as the product (i.e. multiplication) of the row total and the column total, divided by the total number of observations:¹⁰

$$E_{ij} = \frac{R_i \times C_j}{N}$$

Now that we've figured out how to calculate the expected frequencies, it's straightforward to define a test statistic; following the exact same strategy that we used in the goodness of fit test. In fact, it's pretty much the *same* statistic. For a contingency table with r rows and c columns, the equation that defines our X^2 statistic is

$$X^2 = \sum_{i=1}^r \sum_{j=1}^c \frac{(E_{ij} - O_{ij})^2}{E_{ij}}$$

The only difference is that I have to include two summation sign (i.e., \sum) to indicate that we're summing over both rows and columns. As before, large values of X^2 indicate that the null hypothesis provides a poor description of the data, whereas small values of X^2 suggest that it does a good job of accounting for the data. Therefore, just like last time, we want to reject the null hypothesis if X^2 is too large.

Not surprisingly, this statistic is χ^2 distributed. All we need to do is figure out how many degrees of freedom are involved, which actually isn't too hard. As I mentioned before, you can (usually) think of the degrees of freedom as being equal to the number of data points that you're analysing, minus the number of constraints. A contingency table with r rows and c columns contains a total of $r \times c$ observed frequencies, so that's the total number of observations. What about the constraints? Here, it's slightly trickier. The answer is always the same

$$df = (r - 1)(c - 1)$$

but the explanation for *why* the degrees of freedom takes this value is different depending on the experimental design. For the sake of argument, let's suppose that we had honestly intended to survey exactly 87 robots and 93 humans (column totals fixed by the experimenter), but left the row totals free to vary (row totals are random variables). Let's think about the constraints that apply here. Well, since we deliberately fixed the column totals by Act of Experimenter, we have c constraints right there. But, there's actually more to it than that. Remember how our null hypothesis had some free parameters (i.e., we had to estimate the P_i values)? Those matter too. I won't explain why in this book, but every free parameter in the null hypothesis is rather like an additional constraint. So, how many of those are there? Well, since these probabilities have to sum to 1, there's only $r - 1$ of these. So our total degrees of freedom is:

$$\begin{aligned} df &= (\text{number of observations}) - (\text{number of constraints}) \\ &= (rc) - (c + (r - 1)) \\ &= rc - c - r + 1 \\ &= (r - 1)(c - 1) \end{aligned}$$

Alternatively, suppose that the only thing that the experimenter fixed was the total sample size N . That is, we quizzed the first 180 people that we saw, and it just turned out that 87 were robots and 93 were humans. This time around our reasoning would be slightly different, but would still lead us to the same answer. Our null hypothesis still has $r - 1$ free parameters corresponding to the choice probabilities, but it now *also* has $c - 1$ free parameters corresponding to the species probabilities, because we'd also have to estimate the probability that a randomly sampled person turns out to be a robot.¹¹ Finally, since we did actually fix the total number of observations N , that's one more constraint. So now we have, rc

¹⁰Technically, E_{ij} here is an estimate, so I should probably write it \hat{E}_{ij} . But since no-one else does, I won't either.

¹¹A problem many of us worry about in real life.

observations, and $(c - 1) + (r - 1) + 1$ constraints. What does that give?

$$\begin{aligned} df &= (\text{number of observations}) - (\text{number of constraints}) \\ &= rc - ((c - 1) + (r - 1) + 1) \\ &= rc - c - r + 1 \\ &= (r - 1)(c - 1) \end{aligned}$$

Amazing.

12.2.2 Doing the test in R

Okay, now that we know how the test works, let's have a look at how it's done in R. As tempting as it is to lead you through the tedious calculations so that you're forced to learn it the long way, I figure there's no point. I already showed you how to do it the long way for the goodness of fit test in the last section, and since the test of independence isn't conceptually any different, you won't learn anything new by doing it the long way. So instead, I'll go straight to showing you the easy way, which is to use the `chisq.test()` function again. One nice thing about the `chisq.test()` function is that if the `x` argument is a contingency table (rather than a vector), and you don't specify the `p` argument at all, it assumes that you want to run the independence test that I've just described (which I've seen variously referred to as a test of independence, association, homogeneity or contingency). Regardless of what it's called, it's the chi-square test that Pearson developed; and here's how you run it:

```
> chisq.test( x = chapek9.xtabs )  
  
Pearson's Chi-squared test  
  
data: chapek9.xtabs  
X-squared = 10.7216, df = 2, p-value = 0.004697
```

Assuming that descriptive statistics had been provided earlier, we could write up the result.

Pearson's χ^2 revealed a significant association between species and choice ($\chi^2(2) = 10.7, p < .01$): robots appeared to be more likely to say that they prefer flowers, but the humans were more likely to say they prefer data.

Notice that, once again, I provided a little bit of interpretation to help the human reader understand what's going on with the data. Later on in my discussion section, I'd provide a bit more context. To illustrate the difference, here's what I'd probably say later on:

The fact that humans appeared to have a stronger preference for raw data files than robots is somewhat counterintuitive. However, in context it makes some sense: the civil authority on Chapek 9 has an unfortunate tendency to kill and dissect humans when they are identified. As such it seems most likely that the human participants did not respond honestly to the question, so as to avoid potentially undesirable consequences. This should be considered to be a substantial methodological weakness.

This could be classified as a rather extreme example of a reactivity effect, I suppose. Obviously, in this case the problem is severe enough that the study is more or less worthless as a tool for understanding the difference preferences among humans and robots. However, I hope this illustrates the difference between getting a statistically significant result (our null hypothesis is rejected in favour of the alternative), and finding something of scientific value (the data tell us nothing of interest about our research hypothesis due to a big methodological flaw).

12.2.3 Postscript

I later found out the data were made up, and I'd been watching cartoons instead of doing work.

12.3

The continuity correction

Okay, time for a little bit of a digression. I've been lying to you a little bit so far. There's a tiny change that you need to make to your calculations whenever you only have 1 degree of freedom. It's called the "continuity correction", or sometimes the **Yates correction**. Remember what I pointed out earlier: the χ^2 test is based on an approximation, specifically on the assumption that binomial distribution starts to look like a normal distribution for large N . One problem with this is that it often doesn't quite work, especially when you've only got 1 degree of freedom (e.g., when you're doing a test of independence on a 2×2 contingency table). The main reason for this is that the true sampling distribution for the X^2 statistic is actually discrete (because you're dealing with categorical data!) but the χ^2 distribution is continuous. This can introduce systematic problems. Specifically, when N is small and when $df = 1$, the goodness of fit statistic tends to be "too big", meaning that you actually have a bigger α value than you think (or, equivalently, the p values are a bit too small). [Yates \(1934\)](#) suggested a simple fix, in which you redefine the goodness of fit statistic as:

$$X^2 = \sum_i \frac{(|E_i - O_i| - 0.5)^2}{E_i} \tag{12.1}$$

Basically, he just subtracts off 0.5 everywhere. As far as I can tell from reading Yates' paper, the correction is basically a hack. It's not derived from any principled theory: rather, it's based on an examination of the behaviour of the test, and observing that the corrected version seems to work better. I feel obliged to explain this because you will sometimes see R (or any other software for that matter) introduce this correction, so it's kind of useful to know what they're about.

12.4

Effect size

As we discussed earlier (Section 11.8), it's becoming commonplace to ask researchers to report some measure of effect size. So, let's suppose that you've run your chi-square test, which turns out to be significant. So you now know that there is some association between your variables (independence test) or some deviation from the specified probabilities (goodness of fit test). Now you want to report a measure of effect size. That is, given that there is an association/deviation, how strong is it?

There are several different measures that you can choose to report, and several different tools that you can use to calculate them. I won't discuss all of them,¹² but will instead focus on the most commonly reported measures of effect size.

By default, the two measures that people tend to report most frequently are the ϕ statistic and the somewhat superior version, known as Cramér's V . Mathematically, they're very simple. To calculate the

¹²Though I do feel that it's worth mentioning the `assocstats()` function in the `vcd` package. If you install and load the `vcd` package, then a command like `assocstats(chapek9.xtabs)` will run the χ^2 test as well as the likelihood ratio test (not discussed here); and then report three different measures of effect size: ϕ^2 , Cramér's V , and the contingency coefficient (not discussed here)

ϕ statistic, you just divide your X^2 value by the sample size, and take the square root:

$$\phi = \sqrt{\frac{X^2}{N}}$$

The idea here is that the ϕ statistic is supposed to range between 0 (no at all association) and 1 (perfect association), but it doesn't always do this when your contingency table is bigger than 2×2 , which is a total pain. For bigger tables it's actually possible to obtain $\phi > 1$, which is pretty unsatisfactory. So, to correct for this, people usually prefer to report the V statistic proposed by Cramér (1946). It's a pretty simple adjustment to ϕ . If you've got a contingency table with r rows and c columns, then define $k = \min(r, c)$ to be the smaller of the two values. If so, then Cramér's V statistic is

$$V = \sqrt{\frac{X^2}{N(k - 1)}}$$

And you're done. This seems to be a fairly popular measure, presumably because it's easy to calculate, and it gives answers that aren't completely silly: you know that V really does range from 0 (no at all association) to 1 (perfect association).

Calculating V or ϕ is obviously pretty straightforward. So much so that the core packages in R don't seem to have functions to do it, though other packages do. To save you the time and effort of finding one, I've included one in the `lsr` package, called `cramersV()`. For the Chapek 9 data, we get

```
> cramersV( chapek9.xtabs )
[1] 0.244058
```

12.5

Assumptions of the test(s)

All statistical tests make assumptions, and it's usually a good idea to check that those assumptions are met. For the chi-square tests discussed so far in this chapter, the assumptions are:

- *Expected frequencies are sufficiently large.* Remember how in the previous section we saw that the χ^2 sampling distribution emerges because the binomial distribution is pretty similar to a normal distribution? Well, like we discussed in Chapter 9 this is only true when the number of observations is sufficiently large. What that means in practice is that all of the expected frequencies need to be reasonably big. How big is reasonably big? Opinions differ, but the default assumption seems to be that you generally would like to see all your expected frequencies larger than about 5, though for larger tables you would probably be okay if at least 80% of the the expected frequencies are above 5 and none of them are below 1. However, from what I've been able to discover (e.g., Cochran, 1954), these seem to have been proposed as rough guidelines, not hard and fast rules; and they seem to be somewhat conservative (Larntz, 1978).
- *Data are independent of one another.* One somewhat hidden assumption of the chi-square test is that you have to genuinely believe that the observations are independent. Here's what I mean. Suppose I'm interested in proportion of babies born at a particular hospital that are boys. I walk around the maternity wards, and observe 20 girls and only 10 boys. Seems like a pretty convincing difference, right? But later on, it turns out that I'd actually walked into the same ward 10 times, and in fact I'd only seen 2 girls and 1 boy. Not as convincing, is it? My original 30 *observations* were massively non-independent... and were only in fact equivalent to 3 independent observations.

Obviously this is an extreme (and extremely silly) example, but it illustrates the basic issue. Non-independence “stuffs things up”. Sometimes it causes you to falsely reject the null, as the silly hospital example illustrates, but it can go the other way too. To give a slightly less stupid example, let’s consider what would happen if I’d done the cards experiment slightly differently: instead of asking 200 people to try to imagine sampling one card at random, suppose I asked 50 people to select 4 cards. One possibility would be that *everyone* selects one heart, one club, one diamond and one spade (in keeping with the “representativeness heuristic”; Tversky & Kahneman 1974). This is highly non-random behaviour from people, but in this case, I would get an observed frequency of 50 four all four suits. For this example, the fact that the observations are non-independent (because the four cards that you pick will be related to each other) actually leads to the opposite effect... falsely retaining the null.

If you happen to find yourself in a situation where independence is violated, it may be possible to use the McNemar test (which we’ll discuss) or the Cochran test (which we won’t). Similarly, if your expected cell counts are too small, check out the Fisher exact test. It is to these topics that we now turn.

12.6

The Fisher exact test

What should you do if your cell counts are too small, but you’d still like to test the null hypothesis that the two variables are independent? One answer would be “collect more data”, but that’s far too glib: there are a lot of situations in which it would be either infeasible or unethical do that. If so, statisticians have a kind of moral obligation to provide scientists with better tests. In this instance, Fisher (1922) kindly provided the right answer to the question. To illustrate the basic idea, let’s suppose that we’re analysing data from a field experiment, looking at the emotional status of people who have been accused of witchcraft; some of whom are currently being burned at the stake.¹³ Unfortunately for the scientist (but rather fortunately for the general populace), it’s actually quite hard to find people in the process of being set on fire, so the cell counts are awfully small in some cases. The `salem.Rdata` file illustrates the point:

```
> load("salem.Rdata")

> who(TRUE)
-- Name --  -- Class --  -- Size --
trial      data.frame   16 x 2
  $happy    logical      16
  $on.fire  logical      16

> salem.tabs <- table( trial )
> print( salem.tabs )
      on.fire
happy  FALSE TRUE
  FALSE     3     3
  TRUE      10    0
```

Looking at this data, you’d be hard pressed not to suspect that people not on fire are more likely to be happy than people on fire. However, the chi-square test makes this very hard to test because of the small sample size. If I try to do so, R gives me a warning message:

¹³This example is based on a joke article published in the *Journal of Irreproducible Results*.

```

> chisq.test( salem.tabs )

Pearson's Chi-squared test with Yates' continuity correction

data: salem.tabs
X-squared = 3.3094, df = 1, p-value = 0.06888

Warning message:
In chisq.test(salem.tabs) : Chi-squared approximation may be incorrect

```

Speaking as someone who doesn't want to be set on fire, I'd *really* like to be able to get a better answer than this. This is where **Fisher's exact test** (Fisher, 1922) comes in very handy.

The Fisher exact test works somewhat differently to the chi-square test (or in fact any of the other hypothesis tests that I talk about in this book) insofar as it doesn't have a test statistic; it calculates the *p*-value "directly". I'll explain the basics of how the test works for a 2×2 contingency table, though the test works fine for larger tables. As before, let's have some notation:

	Happy	Sad	Total
Set on fire	O_{11}	O_{12}	R_1
Not set on fire	O_{21}	O_{22}	R_2
Total	C_1	C_2	N

In order to construct the test Fisher treats both the row and column totals (R_1 , R_2 , C_1 and C_2) are known, fixed quantities; and then calculates the probability that we would have obtained the observed frequencies that we did (O_{11} , O_{12} , O_{21} and O_{22}) given those totals. In the notation that we developed in Chapter 9 this is written:

$$P(O_{11}, O_{12}, O_{21}, O_{22} | R_1, R_2, C_1, C_2)$$

and as you might imagine, it's a slightly tricky exercise to figure out what this probability is, but it turns out that this probability is described by a distribution known as the *hypergeometric distribution*¹⁴. Now that we know this, what we have to do to calculate our *p*-value is calculate the probability of observing this particular table *or a table that is "more extreme"*.¹⁵ Back in the 1920s, computing this sum was daunting even in the simplest of situations, but these days it's pretty easy as long as the tables aren't too big and the sample size isn't too large. The conceptually tricky issue is to figure out what it means to say that one contingency table is more "extreme" than another. The easiest solution is to say that the table with the lowest probability is the more extreme. This then gives us the *p*-value.

The implementation of the test in R is via the `fisher.test()` function. Here's how it is used:

```

> fisher.test( salem.tabs )

Fisher's Exact Test for Count Data

data: salem.tabs
p-value = 0.03571
alternative hypothesis: true odds ratio is not equal to 1
95 percent confidence interval:
0.000000 1.202913
sample estimates:
odds ratio
0

```

¹⁴The R functions for this distribution are `dhyper()`, `phyper()`, `qhyper()` and `rhyper()`, though you don't need them for this book, and I haven't given you enough information to use these to perform the Fisher exact test the long way.

¹⁵Not surprisingly, the Fisher exact test is motivated by Fisher's interpretation of a *p*-value, not Neyman's!

This is a bit more output than we got from some of our earlier tests. The main thing we're interested in here is the p -value, which in this case is small enough ($p = .036$) to justify rejecting the null hypothesis that people on fire are just as happy as people not on fire.

12.7

The McNemar test

Suppose you've been hired to work for the *Australian Generic Political Party* (AGPP), and part of your job is to find out how effective the AGPP political advertisements are. So, what you do, is you put together a sample of $N = 100$ people, and ask them to watch the AGPP ads. Before they see anything, you ask them if they intend to vote for the AGPP; and then after showing the ads, you ask them again, to see if anyone has changed their minds. Obviously, if you're any good at your job, you'd also do a whole lot of other things too, but let's consider just this one simple experiment. One way to describe your data is via the following contingency table:

	Before	After	Total
Yes	30	10	40
No	70	90	160
Total	100	100	200

At first pass, you might think that this situation lends itself to the Pearson χ^2 test of independence (as per Section 12.2). However, a little bit of thought reveals that we've got a problem: we have 100 participants, but 200 observations. This is because each person has provided us with an answer in *both* the before column and the after column. What this means is that the 200 observations aren't independent of each other: if voter A says "yes" the first time and voter B says "no", then you'd expect that voter A is more likely to say "yes" the second time than voter B! The consequence of this is that the usual χ^2 test won't give trustworthy answers due to the violation of the independence assumption. Now, if this were a really uncommon situation, I wouldn't be bothering to waste your time talking about it. But it's not uncommon at all: this is a *standard* repeated measures design, and none of the tests we've considered so far can handle it. Eek.

The solution to the problem was published by [McNemar \(1947\)](#). The trick is to start by tabulating your data in a slightly different way:

	Before: Yes	Before: No	Total
After: Yes	5	5	10
After: No	25	65	90
Total	30	70	100

This is exactly the same data, but it's been rewritten so that each of our 100 participants appears in only one cell. Because we've written our data this way, the independence assumption is now satisfied, and this is a contingency table that we *can* use to construct an X^2 goodness of fit statistic. However, as we'll see, we need to do it in a slightly nonstandard way. To see what's going on, it helps to label the entries in our table a little differently:

	Before: Yes	Before: No	Total
After: Yes	a	b	$a + b$
After: No	c	d	$c + d$
Total	$a + c$	$b + d$	n

Next, let's think about what our null hypothesis is: it's that the "before" test and the "after" test have the same proportion of people saying "Yes, I will vote for AGPP". Because of the way that we have rewritten the data, it means that we're now testing the hypothesis that the *row totals* and *column totals* come from the same distribution. Thus, the null hypothesis in McNemar's test is that we have "marginal homogeneity". That is, the row totals and column totals have the same distribution: $P_a + P_b = P_a + P_c$, and similarly that $P_c + P_d = P_b + P_d$. Notice that this means that the null hypothesis actually simplifies to $P_b = P_c$. In other words, as far as the McNemar test is concerned, it's only the off-diagonal entries in this table (i.e., b and c) that matter! After noticing this, the **McNemar test of marginal homogeneity** is no different to a usual χ^2 test. After applying the Yates correction, our test statistic becomes:

$$X^2 = \frac{(|b - c| - 0.5)^2}{b + c}$$

or, to revert to the notation that we used earlier in this chapter:

$$X^2 = \frac{(|O_{12} - O_{21}| - 0.5)^2}{O_{12} + O_{21}}$$

and this statistic has an (approximately) χ^2 distribution with $df = 1$. However, remember that – just like the other χ^2 tests – it's only an approximation, so you need to have reasonably large expected cell counts for it to work.

12.7.1 Doing the McNemar test in R

Now that you know what the McNemar test is all about, let's actually run one. The `agpp.Rdata` file contains the raw data that I discussed previously, so let's have a look at it:

```
> load("agpp.Rdata")
> who(TRUE)
-- Name --      -- Class --      -- Size --
agpp           data.frame     100 x 3
$id            factor        100
$response_before factor        100
$response_after factor        100
```

The `agpp` data frame contains three variables, an `id` variable that labels each participant in the data set (we'll see why that's useful in a moment), a `response_before` variable that records the person's answer when they were asked the question the first time, and a `response_after` variable that shows the answer that they gave when asked the same question a second time. As usual, here's the first 6 entries:

```
> head(agpp)
   id response_before response_after
1 subj.1       no          yes
2 subj.2      yes         no
3 subj.3      yes         no
4 subj.4      yes         no
5 subj.5       no         no
6 subj.6       no         no
```

and here's a summary:

```
> summary(agpp)
   id      response_before response_after
```

```

subj.1  : 1    no :70          no :90
subj.10 : 1   yes:30        yes:10
subj.100: 1
subj.11 : 1
subj.12 : 1
subj.13 : 1
(Other) :94

```

Notice that each participant appears only once in this data frame. When we tabulate this data frame using `xtabs()`, we get the appropriate table:

```

> right.table <- xtabs(~ response_before + response_after, data = agpp)
> print(right.table)
      response_after
response_before no yes
      no    65   5
      yes   25   5

```

and from there, we can run the McNemar test by using the `mcnemar.test()` function:

```

> mcnemar.test(right.table)

McNemar's Chi-squared test with continuity correction

data: right.table
McNemar's chi-squared = 12.0333, df = 1, p-value = 0.0005226

```

And we're done. We've just run a McNemar's test to determine if people were just as likely to vote AGPP after the ads as they were before hand. The test was significant ($\chi^2(1) = 12.04, p < .001$), suggesting that they were not. And in fact, it looks like the ads had a negative effect: people were less likely to vote AGPP after seeing the ads. Which makes a lot of sense when you consider the quality of a typical political advertisement.

12.8

Summary

12.8.1 Key concepts

- The Pearson chi-square goodness of fit test is used when you have a table of observed frequencies of different categories; and the null hypothesis gives you a set of “known” probabilities to compare them to. The R function for this looks something like this `chisq.test(x = observed.frequencies, p = null.hyp.probabilities)`.
- The Pearson chi-square test of independence is used when you have a contingency table (cross-tabulation) of two categorical variables. The null hypothesis is that there is no relationship/association between the variables. You use the same function to do this test, but your input should be a contingency table: something like `chisq.test(x = contingency.table)`.
- Effect size for a contingency table can be measured in several ways. In particular we noted the Cramér’s *V* statistic, which can be calculated using `cramersV()`.

- Both versions of the Pearson test rely on two assumptions: that the expected frequencies are sufficiently large, and that the observations are independent. The Fisher exact test can be used when the expected frequencies are small, `fisher.test(x = contingency.table)`. The McNemar test can be used for some kinds of violations of independence, `mcnemar.test(x = contingency.table)`.

12.8.2 Where to next

If you're interested in learning more about categorical data analysis, a good first choice would be [Agresti \(1996\)](#) which, as the title suggests, provides an *Introduction to Categorical Data Analysis*. If the introductory book isn't enough for you (or can't solve the problem you're working on) you could consider [Agresti \(2002\)](#), *Categorical Data Analysis*. The latter is a more advanced text, so it's probably not wise to jump straight from this book to that one.

13. Comparing two means

In the previous chapter we covered the situation when your outcome variable is nominal scale and your predictor variable¹ is also nominal scale. Lots of real world situations have that character, and so you'll find that chi-square tests in particular are quite widely used. However, you're much more likely to find yourself in a situation where your outcome variable is interval scale or higher, and what you're interested in is whether the average value of the outcome variable is higher in one group or another. For instance, a psychologist might want to know if anxiety levels are higher among parents than non-parents, or if working memory capacity is reduced by listening to music (relative to not listening to music). In a medical context, we might want to know if a new drug increases or decreases blood pressure. An agricultural scientist might want to know whether adding phosphorus to Australian native plants will kill them.² In all these situations, our outcome variable is a fairly continuous, interval or ratio scale variable; and our predictor is a binary “grouping” variable. In other words, the problem that we want to address is to compare the means of the two groups.

The standard answer to the problem of comparing means is to use a *t*-test, of which there are several varieties depending on exactly what question you want to solve. As a consequence, the majority of this chapter focuses on different types of *t*-test: one sample *t*-tests are discussed in Section 13.2, independent samples *t*-tests are discussed in Sections 13.3 and 13.4, and paired samples *t*-tests are discussed in Section 13.5. After that, we'll talk a bit about Cohen's *d*, which is the standard measure of effect size for a *t*-test (Section 13.6). The later sections of the chapter focus on the assumptions of the *t*-tests, and possible remedies if they are violated. However, before discussing any of these useful things, we'll start with a discussion of the *z*-test.

13.1

The one-sample *z*-test

In this section I'll describe one of the most useless tests in all of statistics: the *z-test*. Seriously – this test is almost never used in real life. Its only real purpose is that, when teaching statistics, it's a very convenient stepping stone along the way towards the *t*-test, which is probably the most (over)used tool in all statistics.

¹We won't cover multiple predictors until Chapter 15

²Informal experimentation in my garden suggests that yes, it does. Australian natives are adapted to low phosphorus levels relative to everywhere else on Earth, apparently, so if you've bought a house with a bunch of exotics and you want to plant natives, don't follow my example: keep them separate. Nutrients to European plants are poison to Australian ones. There's probably a joke in that, but I can't figure out what it is.

13.1.1 The inference problem that the test addresses

To introduce the idea behind the z -test, let's use a simple example. A friend of mine, Dr Zeppo, grades his introductory statistics class on a curve. Let's suppose that the average grade in his class is 67.5, and the standard deviation is 9.5. Of his many hundreds of students, it turns out that 20 of them also take psychology classes. Out of curiosity, I find myself wondering: do the psychology students tend to get the same grades as everyone else (i.e., mean 67.5) or do they tend to score higher or lower? He emails me the `zeppo.Rdata` file, which I use to pull up the `grades` of those students,

```
> load("zeppo.Rdata")
> print(grades)
[1] 50 60 60 64 66 66 67 69 70 74 76 76 77 79 79 79 81 82 82 89
```

and calculate the mean:

```
> mean(grades)
[1] 72.3
```

Hm. It *might* be that the psychology students are scoring a bit higher than normal: that sample mean of $\bar{X} = 72.3$ is a fair bit higher than the hypothesised population mean of $\mu = 67.5$, but on the other hand, a sample size of $N = 20$ isn't all that big. Maybe it's pure chance.

To answer the question, it helps to be able to write down what it is that I think I know. Firstly, I know that the sample mean is $\bar{X} = 72.3$. If I'm willing to assume that the psychology students have the same standard deviation as the rest of the class then I can say that the population standard deviation is $\sigma = 9.5$. I'll also assume that since Dr Zeppo is grading to a curve, the psychology student grades are normally distributed.

Next, it helps to be clear about what I want to learn from the data. In this case, my research hypothesis relates to the *population* mean μ for the psychology student grades, which is unknown. Specifically, I want to know if $\mu = 67.5$ or not. Given that this is what I know, can we devise a hypothesis test to solve our problem? The data, along with the hypothesised distribution from which they are thought to arise, are shown in Figure 13.1. Not entirely obvious what the right answer is, is it? For this, we are going to need some statistics.

13.1.2 Constructing the hypothesis test

The first step in constructing a hypothesis test is to be clear about what the null and alternative hypotheses are. This isn't too hard to do. Our null hypothesis, H_0 , is that the true population mean μ for psychology student grades is 67.5%; and our alternative hypothesis is that the population mean *isn't* 67.5%. If we write this in mathematical notation, these hypotheses become,

$$\begin{aligned} H_0 : \mu &= 67.5 \\ H_1 : \mu &\neq 67.5 \end{aligned}$$

though to be honest this notation doesn't add much to our understanding of the problem, it's just a compact way of writing down what we're trying to learn from the data. The null hypotheses H_0 and the alternative hypothesis H_1 for our test are both illustrated in Figure 13.2. In addition to providing us with these hypotheses, the scenario outlined above provides us with a fair amount of background knowledge that might be useful. Specifically, there are two special pieces of information that we can add:

1. The psychology grades are normally distributed.

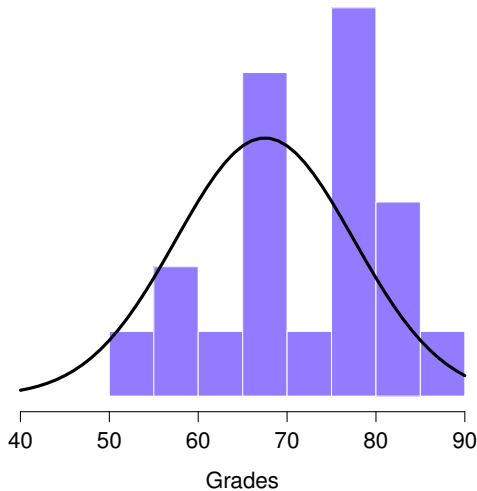


Figure 13.1: The theoretical distribution (solid line) from which the psychology student grades (grey bars) are supposed to have been generated.

2. The true standard deviation of these scores σ is known to be 9.5.

For the moment, we'll act as if these are absolutely trustworthy facts. In real life, this kind of absolutely trustworthy background knowledge doesn't exist, and so if we want to rely on these facts we'll just have make the *assumption* that these things are true. However, since these assumptions may or may not be warranted, we might need to check them. For now though, we'll keep things simple.

The next step is to figure out what we would be a good choice for a diagnostic test statistic; something that would help us discriminate between H_0 and H_1 . Given that the hypotheses all refer to the population mean μ , you'd feel pretty confident that the sample mean \bar{X} would be a pretty useful place to start. What we could do, is look at the difference between the sample mean \bar{X} and the value that the null hypothesis predicts for the population mean. In our example, that would mean we calculate $\bar{X} - 67.5$. More generally, if we let μ_0 refer to the value that the null hypothesis claims is our population mean, then we'd want to calculate

$$\bar{X} - \mu_0$$

If this quantity equals or is very close to 0, things are looking good for the null hypothesis. If this quantity is a long way away from 0, then it's looking less likely that the null hypothesis is worth retaining. But how far away from zero should it be for us to reject H_0 ?

To figure that out, we need to be a bit more sneaky, and we'll need to rely on those two pieces of background knowledge that I wrote down previously, namely that the raw data are normally distributed, and we know the value of the population standard deviation σ . If the null hypothesis is actually true, and the true mean is μ_0 , then these facts together mean that we know the complete population distribution of the data: a normal distribution with mean μ_0 and standard deviation σ . Adopting the notation from Section 9.4, a statistician might write this as:

$$X \sim \text{Normal}(\mu_0, \sigma^2)$$

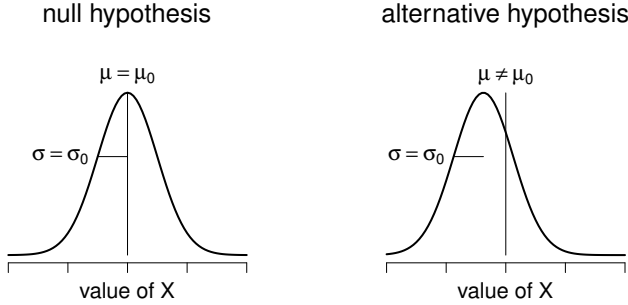


Figure 13.2: Graphical illustration of the null and alternative hypotheses assumed by the one sample z -test (the two sided version, that is). The null and alternative hypotheses both assume that the population distribution is normal, and additionally assumes that the population standard deviation is known (fixed at some value σ_0). The null hypothesis (left) is that the population mean μ is equal to some specified value μ_0 . The alternative hypothesis is that the population mean differs from this value, $\mu \neq \mu_0$.

Okay, if that's true, then what can we say about the distribution of \bar{X} ? Well, as we discussed earlier (see Section 10.4), the sampling distribution of the mean \bar{X} is also normal, and has mean μ . But the standard deviation of this sampling distribution $\text{SE}(\bar{X})$, which is called the *standard error of the mean*, is

$$\text{SE}(\bar{X}) = \frac{\sigma}{\sqrt{N}}$$

In other words, if the null hypothesis is true then the sampling distribution of the mean can be written as follows:

$$\bar{X} \sim \text{Normal}(\mu_0, \text{SE}(\bar{X}))$$

Now comes the trick. What we can do is convert the sample mean \bar{X} into a standard score (Section 5.6). This is conventionally written as z , but for now I'm going to refer to it as $z_{\bar{X}}$. (The reason for using this expanded notation is to help you remember that we're calculating standardised version of a sample mean, *not* a standardised version of a single observation, which is what a z -score usually refers to). When we do so, the z -score for our sample mean is

$$z_{\bar{X}} = \frac{\bar{X} - \mu_0}{\text{SE}(\bar{X})}$$

or, equivalently

$$z_{\bar{X}} = \frac{\bar{X} - \mu_0}{\sigma/\sqrt{N}}$$

This z -score is our test statistic. The nice thing about using this as our test statistics is that like all z -scores, our statistic has a standard normal distribution:

$$z_{\bar{X}} \sim \text{Normal}(0, 1)$$

(again, see Section 5.6 if you've forgotten why this is true). In other words, regardless of what scale the original data are on, the z -statistic itself always has the same interpretation: it's equal to the number of standard errors that separate the observed sample mean \bar{X} from the population mean μ_0 predicted by

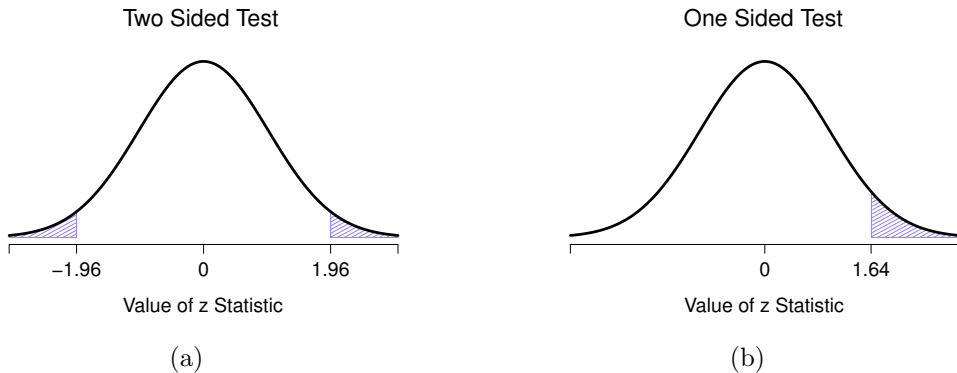


Figure 13.3: Rejection regions for the two-sided z -test (panel a) and the one-sided z -test (panel b).

the null hypothesis. Better yet, regardless of what the population parameters for the raw scores actually are, the 5% critical regions for z -test are always the same, as illustrated in Figure 13.3. And what this meant, way back in the days where people did all their statistics by hand, is that someone could publish a table like this:

desired α level	critical z value	
	two-sided test	one-sided test
.1	1.644854	1.281552
.05	1.959964	1.644854
.01	2.575829	2.326348
.001	3.290527	3.090232

which in turn meant that researchers could calculate their z -statistic by hand, and then look up the critical value in a text book. That was an incredibly handy thing to be able to do back then, but it's kind of unnecessary these days, since it's trivially easy to do it with software like R.

13.1.3 A worked example using R

Now, as I mentioned earlier, the z -test is almost never used in practice. It's so rarely used in real life that the basic installation of R doesn't have a built in function for it. However, the test is so incredibly simple that it's really easy to do one manually. Let's go back to the data from Dr Zeppo's class. Having loaded the `grades` data, the first thing I need to do is calculate the sample mean:

```
> sample.mean <- mean(grades)
> print(sample.mean)
[1] 72.3
```

Then, I create variables corresponding to known population standard deviation ($\sigma = 9.5$), and the value of the population mean that the null hypothesis specifies ($\mu_0 = 67.5$):

```
> mu.null <- 67.5
> sd.true <- 9.5
```

Let's also create a variable for the sample size. We could count up the number of observations ourselves, and type `N <- 20` at the command prompt, but counting is tedious and repetitive. Let's get R to do the tedious repetitive bit by using the `length()` function, which tells us how many elements there are in a vector:

```
> N <- length( grades )
> print( N )
[1] 20
```

Next, let's calculate the (true) standard error of the mean:

```
> sem.true <- sd.true / sqrt(N)
> print(sem.true)
[1] 2.124265
```

And finally, we calculate our z -score:

```
> z.score <- (sample.mean - mu.null) / sem.true
> print( z.score )
[1] 2.259606
```

At this point, we would traditionally look up the value 2.26 in our table of critical values. Our original hypothesis was two-sided (we didn't really have any theory about whether psych students would be better or worse at statistics than other students) so our hypothesis test is two-sided (or two-tailed) also. Looking at the little table that I showed earlier, we can see that 2.26 is bigger than the critical value of 1.96 that would be required to be significant at $\alpha = .05$, but smaller than the value of 2.58 that would be required to be significant at a level of $\alpha = .01$. Therefore, we can conclude that we have a significant effect, which we might write up by saying something like this:

With a mean grade of 73.2 in the sample of psychology students, and assuming a true population standard deviation of 9.5, we can conclude that the psychology students have significantly different statistics scores to the class average ($z = 2.26$, $N = 20$, $p < .05$).

However, what if want an exact p -value? Well, back in the day, the tables of critical values were huge, and so you could look up your actual z -value, and find the smallest value of α for which your data would be significant (which, as discussed earlier, is the very definition of a p -value). However, looking things up in books is tedious, and typing things into computers is awesome. So let's do it using R instead. Now, notice that the α level of a z -test (or any other test, for that matter) defines the total area “under the curve” for the critical region, right? That is, if we set $\alpha = .05$ for a two-sided test, then the critical region is set up such that the area under the curve for the critical region is .05. And, for the z -test, the critical value of 1.96 is chosen that way because the area in the lower tail (i.e., below -1.96) is exactly .025 and the area under the upper tail (i.e., above 1.96) is exactly .025. So, since our observed z -statistic is 2.26, why not calculate the area under the curve below -2.26 or above 2.26? In R we can calculate this using the `pnorm()` function. For the upper tail:

```
> upper.area <- pnorm( q = z.score, lower.tail = FALSE )
> print( upper.area )
[1] 0.01192287
```

The `lower.tail = FALSE` is me telling R to calculate the area under the curve from 2.26 *and upwards*. If I'd told it that `lower.tail = TRUE`, then R would calculate the area from 2.26 *and below*, and it would give me an answer 0.9880771. Alternatively, to calculate the area from -2.26 and below, we get

```

> lower.area <- pnorm( q = -z.score, lower.tail = TRUE )
> print( lower.area )
[1] 0.01192287

```

Thus we get our p -value:

```

> p.value <- lower.area + upper.area
> print( p.value )
[1] 0.02384574

```

13.1.4 How you'd really do the test in R

Now, doing all this is still a great big pain, and it isn't very true to life. On the off chance that you really, really needed to do a z -test in R in real life, you wouldn't do all this thinking and typing. Too much effort. What you'd actually do is the same thing you'd do in every other situation where you want to do a test that isn't in the basic R distribution: use Google. You hunt around online to see if someone else has already written up a package in R that does all this work for you, install the package, and then use the test.

Ah, what the hell. Let's do that. A few minutes on Google revealed to me that some nice person has written an R package called `TeachingDemos` that has a function called `z.test()`. That looks promising. So I install and load the package using the following commands,

```

> install.packages( "TeachingDemos" )
> library( TeachingDemos )

```

(the tedious R output has been omitted), and we're good to go. The `z.test()` function has several arguments, but only three of them are relevant to our toy example. Firstly, we need to specify `x`, a vector containing the raw data (i.e., the `grades`). Secondly, we need to input a number for `mu`, the value of the true mean under the null hypothesis (i.e., 67.5 in this case). Finally, we need to tell R what the population standard deviation `stdev` is (i.e. 9.5). Putting these together, we can use our shiny new `z.test()` function like this:

```

> z.test( x = grades,      # the raw data
+         mu = 67.5,       # the population mean (according to the null)
+         stdev = 9.5,       # the population std dev. (according to the null)
+ )

```

And when we do that, R prints out the answers on screen:

```

One Sample z-test

data:  grades
z = 2.2596, n = 20.000, Std. Dev. = 9.500, Std. Dev.
of the sample mean = 2.124, p-value = 0.02385
alternative hypothesis: true mean is not equal to 67.5
95 percent confidence interval:
 68.13652 76.46348
sample estimates:
mean of grades
72.3

```

If you look at this output closely, you notice that it's telling us that R has done exactly the same thing that we just did manually, and a little bit more to boot. Since R tends to adopt a standardised format for displaying the results of a hypothesis test, it's worth going through it line by line. First, it reminds us that we're doing a `One Sample z-test`, and then goes on to tell us that it's using the raw data from the `grades` vector. Then, there's a long list of numbers, in which R calculates the sample size, the standard error of the mean (i.e., standard deviation of the sample mean), the *z*-score and then finally the *p*-value:

```
z = 2.2596, n = 20.000, Std. Dev. = 9.500, Std. Dev.  
of the sample mean = 2.124, p-value = 0.02385
```

These numbers are the meat of the test. Nevertheless, R is obviously feeling chatty, so it goes onto remind you that these calculations were all based on the assumption that we were doing a two-sided test, in which the alternative hypothesis is that the `true mean is not equal to 67.5`. The next part of the output is our confidence interval for the mean. That is, when R tells you that

```
95 percent confidence interval:  
68.13652 76.46348
```

it's telling you that there's a 95% chance that the true mean is contained by the range 68.14 and 76.46. Finally, it prints out the `sample estimates` (i.e., key quantities that it has estimated from the sample), though in this case there's only the one thing, namely the mean of the `grades` data. The confidence interval and the sample estimates aren't strictly speaking part of the *z*-test, but they're pretty useful things to know.

13.1.5 Assumptions of the *z*-test

As I've said before, all statistical tests make assumptions. Some tests make reasonable assumptions, while other tests do not. The test I've just described – the one sample *z*-test – makes three basic assumptions. These are:

- *Normality*. As usually described, the *z*-test assumes that the true population distribution is normal.³ This is often pretty reasonable, and not only that, it's an assumption that we can check if we feel worried about it (see Section 13.7).
- *Independence*. The second assumption of the test is that the observations in your data set are not correlated with each other, or related to each other in some funny way. This isn't as easy to check statistically: it relies a bit on good experimental design. An obvious (and stupid) example of something that violates this assumption is a data set where you "copy" the same observation over and over again in your data file: so you end up with a massive "sample size", consisting of only one genuine observation. More realistically, you have to ask yourself if it's really plausible to imagine that each observation is a completely random sample from the population that you're interested in. In practice, this assumption is never met; but we try our best to design studies that minimise the problems of correlated data.
- *Known standard deviation*. The third assumption of the *z*-test is that the true standard deviation of the population is known to the researcher. This is just stupid. In no real world data analysis

³Actually this is too strong. Strictly speaking the *z* test only requires that the sampling distribution of the mean be normally distributed; if the population is normal then it necessarily follows that the sampling distribution of the mean is also normal. However, as we saw when talking about the central limit theorem, it's quite possible (even commonplace) for the sampling distribution to be normal even if the population distribution itself is non-normal. However, in light of the sheer ridiculousness of the assumption that the true standard deviation is known, there really isn't much point in going into details on this front!

problem do you know the standard deviation σ of some population, but are completely ignorant about the mean μ . In other words, this assumption is *always* wrong.

In view of the stupidity of assuming that σ is known, let's see if we can live without it. This takes us out of the dreary domain of the z -test, and into the magical kingdom of the t -test, with unicorns and fairies and leprechauns, and um...

13.2

The one-sample t -test

After some thought, I decided that it might not be safe to assume that the psychology student grades necessarily have the same standard deviation as the other students in Dr Zeppo's class. After all, if I'm entertaining the hypothesis that they don't have the same mean, then why should I believe that they absolutely have the same standard deviation? In view of this, I should really stop assuming that I know the true value of σ . This violates the assumptions of my z -test, so in one sense I'm back to square one. However, it's not like I'm completely bereft of options. After all, I've still got my raw data, and those raw data give me an *estimate* of the population standard deviation:

```
> sd( grades )
[1] 9.520615
```

In other words, while I can't say that I know that $\sigma = 9.5$, I *can* say that $\hat{\sigma} = 9.52$.

Okay, cool. The obvious thing that you might think to do is run a z -test, but using the estimated standard deviation of 9.52 instead of relying on my assumption that the true standard deviation is 9.5. So, we could just type this new number into R and out would come the answer. And you probably wouldn't be surprised to hear that this would still give us a significant result. This approach is close, but it's not *quite* correct. Because we are now relying on an *estimate* of the population standard deviation, we need to make some adjustment for the fact that we have some uncertainty about what the true population standard deviation actually is. Maybe our data are just a fluke ... maybe the true population standard deviation is 11, for instance. But if that were actually true, and we ran the z -test assuming $\sigma = 11$, then the result would end up being *non-significant*. That's a problem, and it's one we're going to have to address.

13.2.1 Introducing the t -test

This ambiguity is annoying, and it was resolved in 1908 by a guy called William Sealy Gosset ([Student, 1908](#)), who was working as a chemist for the Guinness brewery at the time (see [Box, 1987](#)). Because Guinness took a dim view of its employees publishing statistical analysis (they felt it was a trade secret), he published the work under the pseudonym "A Student", and to this day, the full name of the t -test is actually **Student's t -test**. The key thing that Gosset figured out is how we should accommodate the fact that we aren't completely sure what the true standard deviation is.⁴ The answer is that it subtly changes the sampling distribution. In the t -test, our test statistic (now called a t -statistic) is calculated in exactly the same way I mentioned above. If our null hypothesis is that the true mean is μ , but our

⁴Well, sort of. As I understand the history, Gosset only provided a partial solution: the general solution to the problem was provided by Sir Ronald Fisher.

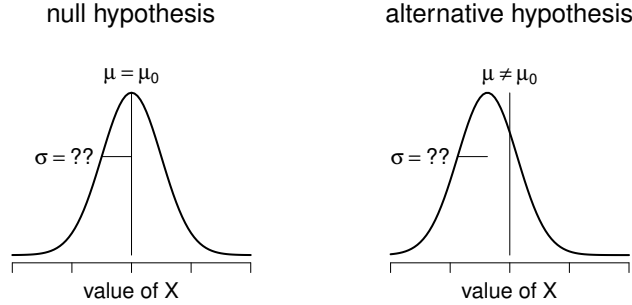


Figure 13.4: Graphical illustration of the null and alternative hypotheses assumed by the (two sided) one sample t -test. Note the similarity to the z -test (Figure 13.2). The null hypothesis is that the population mean μ is equal to some specified value μ_0 , and the alternative hypothesis is that it is not. Like the z -test, we assume that the data are normally distributed; but we do not assume that the population standard deviation σ is known in advance.

sample has mean \bar{X} and our estimate of the population standard deviation is $\hat{\sigma}$, then our t statistic is:

$$t = \frac{\bar{X} - \mu}{\hat{\sigma}/\sqrt{N}}$$

The only thing that has changed in the equation is that instead of using the known true value σ , we use the estimate $\hat{\sigma}$. And if this estimate has been constructed from N observations, then the sampling distribution turns into a t -distribution with $N - 1$ **degrees of freedom** (df). The t distribution is very similar to the normal distribution, but has “heavier” tails, as discussed earlier in Section 9.5 and illustrated in Figure 13.5. Notice, though, that as df gets larger, the t -distribution starts to look identical to the standard normal distribution. This is as it should be: if you have a sample size of $N = 70,000,000$ then your “estimate” of the standard deviation would be pretty much perfect, right? So, you should expect that for large N , the t -test would behave exactly the same way as a z -test. And that’s exactly what happens!

13.2.2 Doing the test in R

As you might expect, the mechanics of the t -test are almost identical to the mechanics of the z -test. So there’s not much point in going through the tedious exercise of showing you how to do the calculations using low level commands: it’s pretty much identical to the calculations that we did earlier, except that we use the estimated standard deviation (i.e., something like `se.est <- sd(grades)`), and then we test our hypothesis using the t distribution rather than the normal distribution (i.e. we use `pt()` rather than `pnorm()`). And so instead of going through the calculations in tedious detail for a second time, I’ll jump straight to showing you how t -tests are actually done in practice. To that end, R provides a `t.test()` function, which is a quite flexible tool. Remember how in Chapter 12 we saw that the `chisq.test()` function can be used to run two different versions of the Pearson chi-square test (the goodness-of-fit test and the test of independence)? Well, the `t.test()` function is even more flexible. There are four different versions of the t -test that it can be used for, all of which we’ll talk about in this chapter. With this in mind, I won’t try to describe all of the arguments to the `t.test()` function now: just the ones that are relevant to the one-sample t -test. They’re pretty similar to the arguments I talked about with the

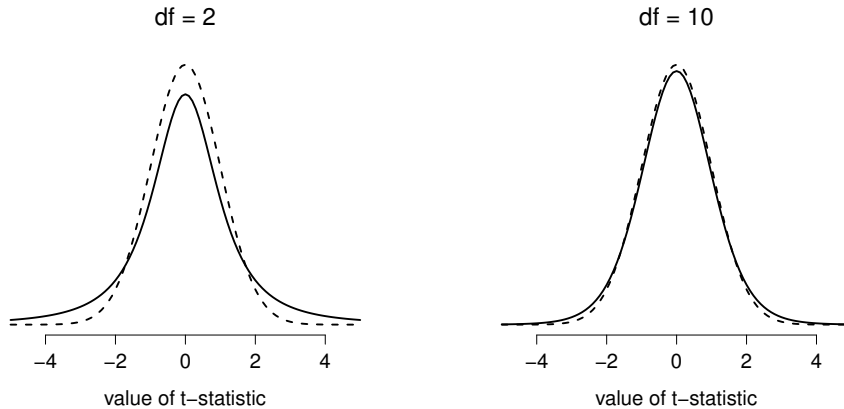


Figure 13.5: The t distribution with 2 degrees of freedom (left) and 10 degrees of freedom (right), with a standard normal distribution (i.e., mean 0 and std dev 1) plotted as dotted lines for comparison purposes. Notice that the t distribution has heavier tails (higher kurtosis) than the normal distribution; this effect is quite exaggerated when the degrees of freedom are very small, but negligible for larger values. In other words, for large df the t distribution is essentially identical to a normal distribution.

`z.test()` function earlier, but I'll cover them again here:

- `x`. A vector containing the sample data.
- `mu`. The population mean that the null hypothesis specifies. If you don't specify a value for `mu`, R will run the test against a value of `mu = 0`.
- `alternative`. This is a character string telling R what type of test to do. There are three possibilities. If you type `alternative = "two.sided"`, R will run a two-sided test. If you say `alternative = "greater"` then R runs a one-sided test in which the null hypothesis is that the mean is (say) 13 or lower, and the alternative hypothesis is that the mean is greater than 13. If you say `alternative = "less"`, then R does the other one-sided test. If you don't specify a value, then R will assume you want a two sided test.
- `conf.level`. The `t.test()` function will also calculate confidence intervals for you, and you can tell it what level of confidence to use. By default, R assumes you want 95% confidence intervals (`conf.level = .95`), which is usually what you want anyway so there's rarely much need to change this.

For the test that I want to run, I'm really only interested in whether the psych student grades are different to everyone else's: I don't have any strong theoretical basis to expect them to be better or worse than other students. Therefore, my null hypothesis is that $\mu = 67.5$, and as a consequence I have a *two-sided* alternative hypothesis ($\mu \neq 67.5$). In other words, I want to run a two-sided test, and hence I don't need to change the default setting of the `alternative` argument. Similarly, I'm going to use the standard α level of .05. I think that 95% confidence intervals are pretty appropriate, and I can leave the `conf.level` argument untouched too. This makes my test pretty easy to run: all I need to do is specify the data `x` and the population mean that the null hypothesis specifies `mu`. Thus, I run my t -test by typing in the following command:

```
> t.test( x = grades, # 'grades' is a numeric vector containing the data
+         mu = 67.5    # the population mean is 67.5 if the null is true
+ )
```

And here's the output:

```
One Sample t-test

data: grades
t = 2.2547, df = 19, p-value = 0.03615
alternative hypothesis: true mean is not equal to 67.5
95 percent confidence interval:
 67.84422 76.75578
sample estimates:
mean of x
 72.3
```

That's much less typing, and R is kind enough to print out everything in a nice, short summary. You'll notice that this has the exact same format as the results from the *z*-test from the previous section. As before, the key numbers are all in this line:

```
t = 2.2547, df = 19, p-value = 0.03615
```

The rest of it is just R being polite and reminding you that it did exactly what you asked, or printing out a few other things that it thinks you'll need (e.g., the confidence interval for the mean).

In any case, since we're pretending that we actually care about my toy example, we're overjoyed to discover that the result is statistically significant. We could report the result by saying something like this:

With a mean grade of 72.3, the psychology students scored slightly higher than the average grade of 67.5 ($t(19) = 2.25, p < .05$); the 95% confidence interval is [67.8, 76.8].

where $t(19)$ is shorthand notation for a *t*-statistic that has 19 degrees of freedom. That said, it's often the case that people don't report the confidence interval, or do so using a much more compressed form than I've done here. For instance, it's not uncommon to see the confidence interval included as part of the stat block, like this:

$t(19) = 2.25, p < .05, \text{CI}_{95} = [67.8, 76.8]$

With that much jargon crammed into half a line, you know it must be really smart.⁵

13.2.3 Assumptions of the one sample *t*-test

Okay, so what assumptions does the one-sample *t*-test make? Well, since the *t*-test is basically a *z*-test with the assumption of known standard deviation removed, you shouldn't be surprised to see that it makes the same assumptions as the *z*-test, minus the one about the known standard deviation. That is

⁵More seriously, I tend to think the reverse is true: I get very suspicious of technical reports that fill their results sections with nothing except the numbers. It might just be that I'm an arrogant jerk, but I often feel like an author that makes no attempt to explain and interpret their analysis to the reader either doesn't understand it themselves, or is being a bit lazy. Your readers are smart, but not infinitely patient. Don't annoy them if you can help it.

- *Normality*. We're still assuming that the population distribution is normal⁶, and as noted earlier, there are standard tools that you can use to check to see if this assumption is met (Section 13.7), and other tests you can do in its place if this assumption is violated (Section 13.8).
- *Independence*. Once again, we have to assume that the observations in our sample are generated independently of one another. See the earlier discussion about the *z*-test for specifics (Section 13.1.5).

Overall, these two assumptions aren't terribly unreasonable, and as a consequence the one-sample *t*-test is pretty widely used in practice as a way of comparing a sample mean against a hypothesised population mean.

13.3

The independent samples *t*-test (Student test)

Although the one sample *t*-test has its uses, it's not the most typical example of a *t*-test⁷. A much more common situation arises when you've got two different groups of observations. In psychology, this tends to correspond to two different groups of participants, where each group corresponds to a different condition in your study. For each person in the study, you measure some outcome variable of interest, and the research question that you're asking is whether or not the two groups have the same population mean. This is the situation that the independent samples *t*-test is designed for.

13.3.1 The data

Suppose we have 33 students taking Dr Harpo's statistics lectures, and Dr Harpo doesn't grade to a curve. Actually, Dr Harpo's grading is a bit of a mystery, so we don't really know anything about what the average grade is for the class as a whole. There are two tutors for the class, Anastasia and Bernadette. There are $N_1 = 15$ students in Anastasia's tutorials, and $N_2 = 18$ in Bernadette's tutorials. The research question I'm interested in is whether Anastasia or Bernadette is a better tutor, or if it doesn't make much of a difference. Dr Harpo emails me the course grades, in the `harpo.Rdata` file. As usual, I'll load the file and have a look at what variables it contains:

```
> load("harpo.Rdata")
> who(TRUE)
-- Name -- -- Class -- -- Size --
harpo      data.frame   33 x 2
$grade     numeric       33
$tutor     factor        33
```

As we can see, there's a single data frame with two variables, `grade` and `tutor`. The `grade` variable is a numeric vector, containing the grades for all $N = 33$ students taking Dr Harpo's class; the `tutor` variable is a factor that indicates who each student's tutor was. The first six observations in this data set are shown below:

⁶A technical comment... in the same way that we can weaken the assumptions of the *z*-test so that we're only talking about the sampling distribution, we *can* weaken the *t* test assumptions so that we don't have to assume normality of the population. However, for the *t*-test, it's trickier to do this. As before, we can replace the assumption of population normality with an assumption that the sampling distribution of \bar{X} is normal. However, remember that we're also relying on a sample estimate of the standard deviation; and so we also require the sampling distribution of $\hat{\sigma}$ to be chi-square. That makes things nastier, and this version is rarely used in practice: fortunately, if the population is normal, then both of these two assumptions are met.

⁷Although it is the simplest, which is why I started with it.

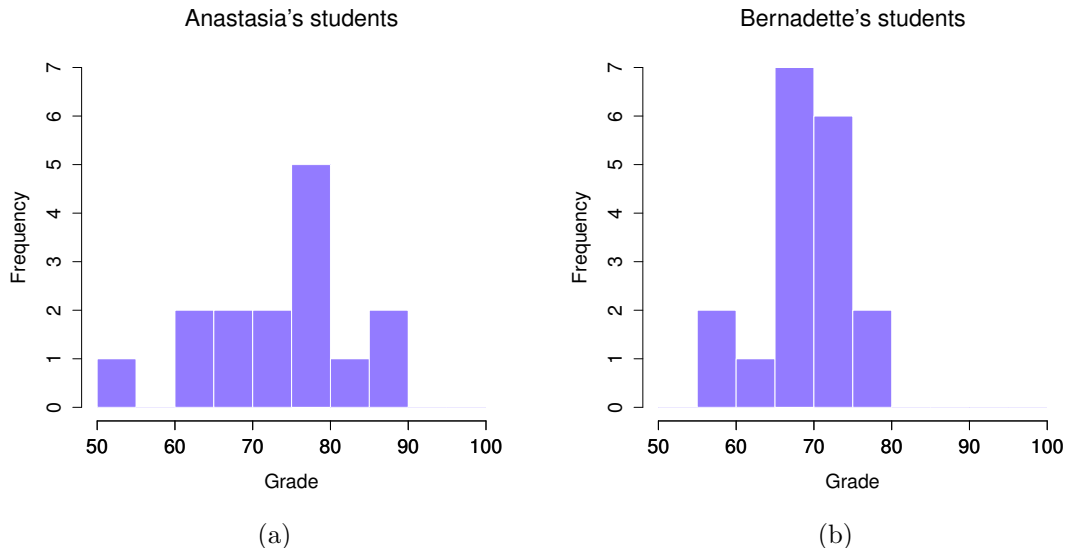


Figure 13.6: Histograms showing the overall distribution of grades for students in Anastasia’s class (panel a) and in Bernadette’s class (panel b). Inspection of these histograms suggests that the students in Anastasia’s class may be getting slightly better grades on average, though they also seem a little more variable.

```
> head( harpo )
  grade      tutor
1    65  Anastasia
2    72  Bernadette
3    66  Bernadette
4    74  Anastasia
5    73  Anastasia
6    71  Bernadette
```

We can calculate means and standard deviations, using the `mean()` and `sd()` functions. Rather than show the R output, here’s a nice little summary table:

	mean	std dev	N
Anastasia’s students	74.53	9.00	15
Bernadette’s students	69.06	5.77	18

To give you a more detailed sense of what’s going on here, I’ve plotted histograms showing the distribution of grades for both tutors (Figure 13.6), as well as a simpler plot showing the means and corresponding confidence intervals for both groups of students (Figure 13.7).

13.3.2 Reorganising the data

Before going on to talk about the independent samples *t*-test itself, I’m going to reorganise the data a little bit. At the moment we’ve got all the data stored in a single data frame. With the data in this format, we can think of `grade` as our outcome variable, and `tutor` as the predictor (also called a

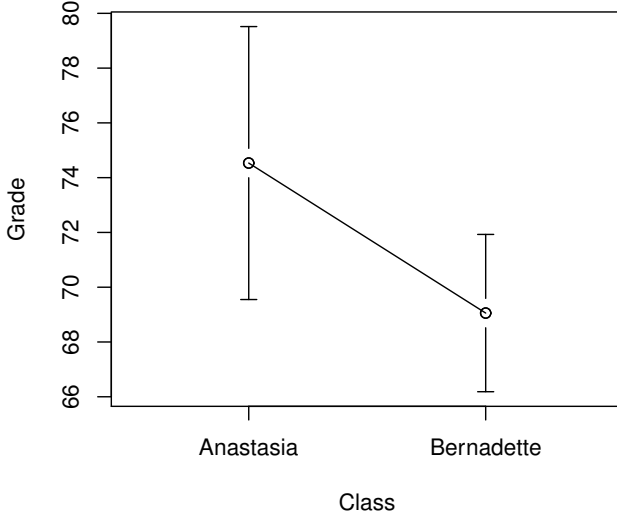


Figure 13.7: Plots showing the mean grade for the students in Anastasia's and Bernadette's tutorials. Error bars depict 95% confidence intervals around the mean. On the basis of visual inspection, it does look like there's a real difference between the groups, though it's hard to say for sure.

grouping variable in this context). This format is probably the way you're most likely to have your data organised in real life, but it's not the only one. As an alternative, we could have two separate variables containing the grades for Anastasia's students `grade.A` and the grades for Bernadette's students `grade.B`. When stored in this format, our variables would look like this:

```
> # Anastasia's students' grades:
> print( grade.A )
[1] 55 65 65 68 70 73 74 76 77 78 78 80 83 86 90
>
> # Bernadette's students' grades:
> print( grade.B )
[1] 56 60 62 66 66 67 68 68 69 69 71 72 73 73 74 74 76 79
```

Thus, we have now two different ways of organising the data: one using `grade` and `tutor` as the variables, and the other using the `grade.A` and `grade.B` variables. When doing an independent samples *t*-test, you might find yourself with data in either of these two formats. As we'll see, however, the `t.test()` function is flexible enough to support both. In case you want to try playing with the data in both formats, here's the commands I used to create the `grade.A` and `grade.B` variables:

```
> grade.A <- with( harpo, grade[tutor == "Anastasia"] )
> grade.B <- with( harpo, grade[tutor == "Bernadette"] )
```

... because I know you're all just so *excited* at the prospect of doing *t*-tests that you'd relish the opportunity to do the same one twice.⁸ So much so that I saved these new variables in the file `harpo2.Rdata`, so you can play with them yourself,

```
> load( "harpo2.Rdata" )
```

13.3.3 Introducing the test

The **independent samples *t*-test** comes in two different forms, Student's and Welch's. The original Student *t*-test – which is the one I'll describe in this section – is the simpler of the two, but relies on much more restrictive assumptions than the Welch *t*-test. Assuming for the moment that you want to run a two-sided test, the goal is to determine whether two “independent samples” of data are drawn from populations with the same mean (the null hypothesis) or different means (the alternative hypothesis). When we say “independent” samples, what we really mean here is that there's no special relationship between observations in the two samples. This probably doesn't make a lot of sense right now, but it will be clearer when we come to talk about the paired samples *t*-test later on. For now, let's just point out that if we have an experimental design where participants are randomly allocated to one of two groups, and we want to compare the two groups' mean performance on some outcome measure, then an independent samples *t*-test (rather than a paired samples *t*-test) is what we're after.

Okay, so let's let μ_1 denote the true population mean for group 1 (e.g., Anastasia's students), and μ_2 will be the true population mean for group 2 (e.g., Bernadette's students),⁹ and as usual we'll let \bar{X}_1 and \bar{X}_2 denote the observed sample means for both of these groups. Our null hypothesis states that the two population means are identical ($\mu_1 = \mu_2$) and the alternative to this is that they are not ($\mu_1 \neq \mu_2$). Written in mathematical-ese, this is...

$$\begin{aligned} H_0 : \mu_1 &= \mu_2 \\ H_1 : \mu_1 &\neq \mu_2 \end{aligned}$$

To construct a hypothesis test that handles this scenario, we start by noting that if the null hypothesis is true, then the difference between the population means is *exactly* zero, $\mu_1 - \mu_2 = 0$. As a consequence, a diagnostic test statistic will be based on the difference between the two sample means. Because if the null hypothesis is true, then we'd expect $\bar{X}_1 - \bar{X}_2$ to be *pretty close* to zero. However, just like we saw with our one-sample tests (i.e., the one-sample *z*-test and the one-sample *t*-test) we have to be precise about exactly *how close* to zero this difference should be. And the solution to the problem is more or less the same one: we calculate a standard error estimate (SE), just like last time, and then divide the difference between means by this estimate. So our ***t*-statistic** will be of the form

$$t = \frac{\bar{X}_1 - \bar{X}_2}{\text{SE}}$$

⁸A similar result can be obtained by using the command `importList(split(harpo$grade, harpo$tutor))`, which will create two new variables called `Anastasia` and `Bernadette`, each of which contains the grades from the relevant class. The `importList` function is in the `lsr` package.

⁹A funny question almost always pops up at this point: what the heck *is* the population being referred to in this case? Is it the set of students actually taking Dr Harpo's class (all 33 of them)? The set of people who might take the class (an unknown number) of them? Or something else? Does it matter which of these we pick? It's traditional in an introductory behavioural stats class to mumble a lot at this point, but since I get asked this question every year by my students, I'll give a brief answer. Technically yes, it does matter: if you change your definition of what the “real world” population actually is, then the sampling distribution of your observed mean \bar{X} changes too. The *t*-test relies on an assumption that the observations are sampled at random from an infinitely large population; and to the extent that real life isn't like that, then the *t*-test can be wrong. In practice, however, this isn't usually a big deal: even though the assumption is almost always wrong, it doesn't lead to a lot of pathological behaviour from the test, so we tend to just ignore it.

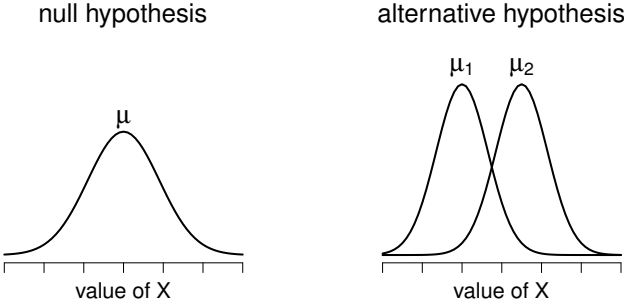


Figure 13.8: Graphical illustration of the null and alternative hypotheses assumed by the Student *t*-test. The null hypothesis assumes that both groups have the same mean μ , whereas the alternative assumes that they have different means μ_1 and μ_2 . Notice that it is assumed that the population distributions are normal, and that, although the alternative hypothesis allows the group to have different means, it assumes they have the same standard deviation.

We just need to figure out what this standard error estimate actually is. This is a bit trickier than was the case for either of the two tests we've looked at so far, so we need to go through it a lot more carefully to understand how it works.

13.3.4 A “pooled estimate” of the standard deviation

In the original “Student *t*-test”, we make the assumption that the two groups have the same population standard deviation: that is, regardless of whether the population means are the same, we assume that the population standard deviations are identical, $\sigma_1 = \sigma_2$. Since we’re assuming that the two standard deviations are the same, we drop the subscripts and refer to both of them as σ . How should we estimate this? How should we construct a single estimate of a standard deviation when we have two samples? The answer is, basically, we average them. Well, sort of. Actually, what we do is take a *weighed* average of the *variance* estimates, which we use as our **pooled estimate of the variance**. The weight assigned to each sample is equal to the number of observations in that sample, minus 1. Mathematically, we can write this as

$$\begin{aligned} w_1 &= N_1 - 1 \\ w_2 &= N_2 - 1 \end{aligned}$$

Now that we’ve assigned weights to each sample, we calculate the pooled estimate of the variance by taking the weighted average of the two variance estimates, $\hat{\sigma}_1^2$ and $\hat{\sigma}_2^2$

$$\hat{\sigma}_p^2 = \frac{w_1 \hat{\sigma}_1^2 + w_2 \hat{\sigma}_2^2}{w_1 + w_2}$$

Finally, we convert the pooled variance estimate to a pooled standard deviation estimate, by taking the square root. This gives us the following formula for $\hat{\sigma}_p$,

$$\hat{\sigma}_p = \sqrt{\frac{w_1 \hat{\sigma}_1^2 + w_2 \hat{\sigma}_2^2}{w_1 + w_2}}$$

And if you mentally substitute $w_1 = N_1 - 1$ and $w_2 = N_2 - 1$ into this equation you get a very ugly looking formula; a very ugly formula that actually seems to be the “standard” way of describing the pooled

standard deviation estimate. It's not my favourite way of thinking about pooled standard deviations, however.¹⁰

13.3.5 The same pooled estimate, described differently

I prefer to think about it like this. Our data set actually corresponds to a set of N observations, which are sorted into two groups. So let's use the notation X_{ik} to refer to the grade received by the i -th student in the k -th tutorial group: that is, X_{11} is the grade received by the first student in Anastasia's class, X_{21} is her second student, and so on. And we have two separate group means \bar{X}_1 and \bar{X}_2 , which we could "generically" refer to using the notation \bar{X}_k , i.e., the mean grade for the k -th tutorial group. So far, so good. Now, since every single student falls into one of the two tutorials, and so we can describe their deviation from the group mean as the difference

$$X_{ik} - \bar{X}_k$$

So why not just use these deviations (i.e., the extent to which each student's grade differs from the mean grade in their tutorial?) Remember, a variance is just the average of a bunch of squared deviations, so let's do that. Mathematically, we could write it like this:

$$\frac{\sum_{ik} (X_{ik} - \bar{X}_k)^2}{N}$$

where the notation " \sum_{ik} " is a lazy way of saying "calculate a sum by looking at all students in all tutorials", since each " ik " corresponds to one student.¹¹ But, as we saw in Chapter 10, calculating the variance by dividing by N produces a biased estimate of the population variance. And previously, we needed to divide by $N - 1$ to fix this. However, as I mentioned at the time, the reason why this bias exists is because the variance estimate relies on the sample mean; and to the extent that the sample mean isn't equal to the population mean, it can systematically bias our estimate of the variance. But this time we're relying on *two* sample means! Does this mean that we've got more bias? Yes, yes it does. And does this mean we now need to divide by $N - 2$ instead of $N - 1$, in order to calculate our pooled variance estimate? Why, yes...

$$\hat{\sigma}_p^2 = \frac{\sum_{ik} (X_{ik} - \bar{X}_k)^2}{N - 2}$$

Oh, and if you take the square root of this then you get $\hat{\sigma}_p$, the pooled standard deviation estimate. In other words, the pooled standard deviation calculation is nothing special: it's not terribly different to the regular standard deviation calculation.

13.3.6 Completing the test

Regardless of which way you want to think about it, we now have our pooled estimate of the standard deviation. From now on, I'll drop the silly p subscript, and just refer to this estimate as $\hat{\sigma}$. Great. Let's now go back to thinking about the bloody hypothesis test, shall we? Our whole reason for calculating this pooled estimate was that we knew it would be helpful when calculating our *standard error* estimate. But, standard error of *what*? In the one-sample t -test, it was the standard error of the sample mean, $SE(\bar{X})$, and since $SE(\bar{X}) = \sigma/\sqrt{N}$ that's what the denominator of our t -statistic looked like. This time around, however, we have *two* sample means. And what we're interested in, specifically, is the the difference between the two $\bar{X}_1 - \bar{X}_2$. As a consequence, the standard error that we need to divide by is in fact the **standard error of the difference** between means. As long as the two variables really do have the

¹⁰Yes, I have a "favourite" way of thinking about pooled standard deviation estimates. So what?

¹¹A more correct notation will be introduced in Chapter 14.

same standard deviation, then our estimate for the standard error is

$$\text{SE}(\bar{X}_1 - \bar{X}_2) = \frac{\hat{\sigma}}{\sqrt{\frac{1}{N_1} + \frac{1}{N_2}}}$$

and our t -statistic is therefore

$$t = \frac{\bar{X}_1 - \bar{X}_2}{\text{SE}(\bar{X}_1 - \bar{X}_2)}$$

Just as we saw with our one-sample test, the sampling distribution of this t -statistic is a t -distribution (shocking, isn't it?) as long as the null hypothesis is true, and all of the assumptions of the test are met. The degrees of freedom, however, is slightly different. As usual, we can think of the degrees of freedom to be equal to the number of data points minus the number of constraints. In this case, we have N observations (N_1 in sample 1, and N_2 in sample 2), and 2 constraints (the sample means). So the total degrees of freedom for this test are $N - 2$.

13.3.7 Doing the test in R, version 1

As I was saying earlier, one of the four different tests that the `t.test()` function can perform is the Student independent samples t -test. When using the test this way, the three main arguments that you need to care about are `x`, `y` and `var.equal`. In this context, what they mean is this:

- `x`. A vector containing the data for sample 1.
- `y`. A vector containing the data for sample 2.
- `var.equal`. You need to explicitly tell the `t.test()` function to make the assumption that standard deviations are equal, by setting `var.equal = TRUE`. If you don't do this, it will run the Welch test instead of the Student test.

As before, you can use the `alternative` argument to tell R to switch between two sided tests and one sided tests, and the `conf.level` argument can be used to set the α level for the confidence intervals it spits out. But these arguments work exactly the same way regardless of which test you're using, so there's nothing new to add about them. With this in mind, to run our test, here's what we type:

```
> t.test( x = grade.A,      # the grades from Anastasia's students
+         y = grade.B,      # the grades from Bernadette's students
+         var.equal = TRUE    # do a Student test, not a Welch test
+ )
```

and here's the output:

```
Two Sample t-test

data: grade.A and grade.B
t = 2.1154, df = 31, p-value = 0.04253
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 0.1965873 10.7589683
sample estimates:
mean of x mean of y
74.53333 69.05556
```

Let's look at it closely. Some parts of the output are just R telling us exactly what test it did: the two sample t-test (i.e., the Student's independent samples t -test). It also tells us what it used as the

alternative hypothesis, that the true difference in means is not equal to 0. The third thing that it feels the need to remind us of is what it used for the data: `grade.A` and `grade.B`.

None of those things ought to surprise you: those are the specifications of the test that *you gave it*. R is literally telling you stuff you already know. It does this for two reasons: (1) this is R politely hinting that you should actually check that you specified the right test, because it's easy to accidentally type the wrong thing, so it's quietly reminding you of what it thinks you want; and (2) because sometimes you might want to save the output of the `t.test()` function to a variable,¹² and when you're looking at it later on you might have forgotten what the test you did actually looks like. This is quite easy to do:

```
> my.t.test <- t.test( x=grade.A, y=grade.B, var.equal=TRUE )
```

When you do this, R stores the results of the test to a variable called `my.t.test`, which you can `print()` out whenever you like. So if I were to then type `print(my.t.test)`, I'd get the exact same output that I showed earlier.

Getting back to the output itself, just like last time the important line is the one that reads `t = 2.1154, df = 31, p-value = 0.04253` because that's the one that tells us that we found a significant difference between the two groups ($t(31) = 2.12, p = .04$). However, the other lines are quite helpful too. When interpreting the test it's usually helpful to know what the `sample estimates` (i.e. sample means) were, so R prints those out too. Finally, it also provides a confidence interval. However, if you take a moment to think about the part of the output says,

```
95 percent confidence interval:  
 0.1965873 10.7589683
```

it's clear that it's not giving us the confidence interval for the individual population means μ_1 and μ_2 . Instead, what you're looking at is the confidence interval for the *difference* between means, $\mu_1 - \mu_2$.

13.3.8 Doing the test in R, version 2

Okay, so that's how we use the `t.test()` function if the data for each group is stored in a separate variable. But that's not always the case. In fact, you're much more likely to have data stored in a data frame (like `harpo`), which would contain an outcome variable (like `grade`) and a grouping variable (like `tutor`). It would be a pain to have to convert those over every time, so R provides a different method for running the test when you have the data in this format. The test works exactly the same as before, except that instead of using `x` and `y` to specify the data, you use these:

- **formula**. A formula of the format `outcome ~ group`. That is, on the left hand side you type the name of the outcome variable, and on the right hand side you type the name of the grouping variable.
- **data**. The data frame where the variables are stored. If this argument isn't specified, R will look in the workspace to see if the variables are there.

For our example, we type in this:

```
> t.test( formula = grade ~ tutor, # outcome ~ group  
+         data = harpo,           # name of data frame  
+         var.equal = TRUE        # Student test, not Welch  
+ )
```

and we obtain an output that is the same in all substantive details:

¹²Actually, you can do this for *any* of the hypothesis test functions that we've talked about so far.

Two Sample t-test

```
data: grade by tutor
t = 2.1154, df = 31, p-value = 0.04253
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 0.1965873 10.7589683
sample estimates:
mean in group Anastasia mean in group Bernadette
 74.53333           69.05556
```

13.3.9 Positive and negative t values

Before moving on to talk about the assumptions of the t -test, there's one additional point I want to make about the use of t -tests in practice. The first one relates to the sign of the t -statistic (that is, whether it is a positive number or a negative one). One very common worry that students have when they start running their first t -test is that they often end up with negative values for the t -statistic, and don't know how to interpret it. In fact, it's not at all uncommon for two people working independently to end up with R outputs that are almost identical, except that one person has a negative t values and the other one has a positive t value. Assuming that you're running a two-sided test, then the p -values will be identical. On closer inspection, the students will notice that the confidence intervals also have the opposite signs. This is perfectly okay: whenever this happens, what you'll find is that the two versions of the R output arise from slightly different ways of running the t -test. What's happening here is very simple. The t -statistic that R is calculating here is always of the form

$$t = \frac{(\text{mean 1}) - (\text{mean 2})}{(\text{SE})}$$

If "mean 1" is larger than "mean 2" the t statistic will be positive, whereas if "mean 2" is larger then the t statistic will be negative. Similarly, the confidence interval that R reports is the confidence interval for the difference "(mean 1) minus (mean 2)", which will be the reverse of what you'd get if you were calculating the confidence interval for the difference "(mean 2) minus (mean 1)".

Okay, that's pretty straightforward when you think about it, but now consider our t -test comparing Anastasia's class to Bernadette's class. Which one should we call "mean 1" and which one should we call "mean 2". It's arbitrary. However, you really do need to designate one of them as "mean 1" and the other one as "mean 2". The way that R handles this is as follows. If you are calling the `t.test()` function by using the `x` and `y` arguments, then R will use `x` as mean 1, and `y` as mean 2. On the other hand, if you're using the `formula` argument, then R will use the same ordering that you would get using the `sort()` function (see Section 7.6). For instance, if your grouping variable is numeric, then "mean 1" will correspond to the smaller group number. If it is a character vector, then they're ordered alphabetically, with "mean 1" corresponding to the group name that comes earlier in the dictionary. Finally, if the grouping variable is a factor, "mean 1" will correspond to the first factor level (usually alphabetically ordered, but not always).

In practice of course, this doesn't make much of a difference, and you probably won't remember all the details of how R determines which group counts as "mean 1". I never remember. What I usually do instead is check the actual group means. That is, whenever I get a significant t -test result, and I want to figure out which mean is the larger one, I don't bother trying to figure it out by looking at the t -statistic: instead, I just look at the actual group means.

Relatedly, irrespective of what R actually prints out, I usually try to *report* the *t*-statistic in such a way that the numbers match up with the text. Here's what I mean... suppose that what I want to write in my report is "Anastasia's class had higher grades than Bernadette's class". The phrasing here has Anastasia's group coming first, so it makes sense to report the *t*-statistic as if Anastasia's class corresponded to group 1. If so, I would write

Anastasia's class had higher grades than Bernadette's class ($t(31) = 2.1, p = .04$).

(I wouldn't actually underline the word "higher" in real life, I'm just doing it to emphasise the point that "higher" corresponds to positive *t* values). On the other hand, suppose the phrasing I wanted to use has Bernadette's class listed first. If so, it makes more sense to treat her class as group 1, and if so, the write up looks like this:

Bernadette's class had lower grades than Anastasia's class ($t(31) = -2.1, p = .04$).

Because I'm talking about one group having "lower" scores this time around, it is more sensible to use the negative form of the *t*-statistic.¹³

13.3.10 Assumptions of the test

The Student independent samples *t*-test relies on three assumptions, some of which we saw previously in the context of the one sample *t*-test (see Section 13.2.3). They are:

- *Normality*. Like the one-sample *t*-test, it is assumed that the data are normally distributed. Specifically, we assume that both groups are normally distributed. In Section 13.7 we'll discuss how to test for normality, and in Section 13.8 we'll discuss possible solutions.
- *Independence*. Once again, it is assumed that the observations are independently sampled. In the context of the Student test this has two aspects to it. Firstly, we assume that the observations within each sample are independent of one another (exactly the same as for the one-sample test). However, we also assume that there are no cross-sample dependencies. If, for instance, it turns out that you included some participants in both experimental conditions of your study (e.g., by accidentally allowing the same person to sign up to different conditions), then there are some cross sample dependencies that you'd need to take into account.
- *Homogeneity of variance* (also called "homoscedasticity"). The third assumption is that the population standard deviation is the same in both groups. You can test this assumption using the Levene test, which I'll talk about later on in the book (Section 14.7). However, there's a very simple remedy for this assumption, which I'll talk about in the next section.

13.4

The independent samples *t*-test (Welch test)

One of the biggest problems with the Student test is the assumption that both groups have exactly the same variance, which is often not satisfied by real data. We'll talk a little bit about how you can check this assumption later on because it does crop up in a few different places, not just the *t*-test. But right now I'll talk about a different form of the *t*-test (Welch, 1947) that does not rely on this assumption.

¹³Please note that you can't do this for other types of test statistics. It works for *t*-tests, but it wouldn't be meaningful for chi-square tests or indeed for most of the tests I talk about in this book. So don't overgeneralise this advice!

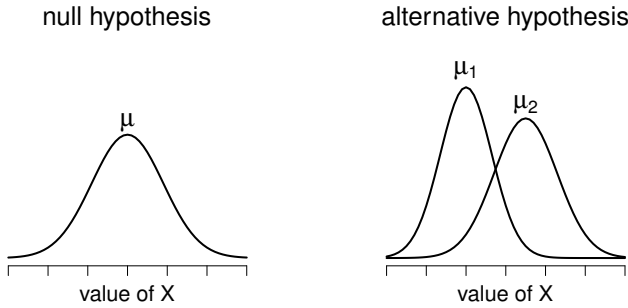


Figure 13.9: Graphical illustration of the null and alternative hypotheses assumed by the Welch t -test. Like the Student test (Figure 13.8) we assume that both samples are drawn from a normal population; but the alternative hypothesis no longer requires the two populations to have equal variance.

.....

A graphical illustration of what the **Welch t test** assumes about the data is shown in Figure 13.9, to provide a contrast with the Student test version in Figure 13.8. I'll admit it's a bit odd to talk about the cure before talking about the diagnosis, but as it happens the Welch test is the default t -test in R, so this is probably the best place to discuss it. I won't go into as much detail this time. The t -statistic that we use in the Welch test has the same basic format as the one used in the Student test, i.e.

$$t = \frac{\bar{X}_1 - \bar{X}_2}{\text{SE}(\bar{X}_1 - \bar{X}_2)}$$

but the standard error calculations are different. If the two populations have different standard deviations, then it's a complete nonsense to try to calculate a pooled standard deviation estimate, because you're averaging apples and oranges.¹⁴ But you can still estimate the standard error of the difference between sample means; it just ends up looking different:

$$\text{SE}(\bar{X}_1 - \bar{X}_2) = \sqrt{\frac{\hat{\sigma}_1^2}{N_1} + \frac{\hat{\sigma}_2^2}{N_2}}$$

What this means is that the t -statistic that comes out of the Welch test is actually somewhat different to the one that comes from the Student test. Worse yet, the degrees of freedom calculations are very very different. The degrees of freedom doesn't have to be a whole number any more, and it certainly doesn't correspond all that closely to the "number of data points minus the number of constraints" heuristic that I've been using up to this point. The degrees of freedom are, in fact...

$$\text{df} = \frac{(\hat{\sigma}_1^2/N_1 + \hat{\sigma}_2^2/N_2)^2}{(\hat{\sigma}_1^2/N_1)^2/(N_1 - 1) + (\hat{\sigma}_2^2/N_2)^2/(N_2 - 1)}$$

... which is all pretty straightforward and obvious, right? Well, perhaps not.

¹⁴Well, I guess you can average apples and oranges, but what you end up with is a delicious fruit smoothie, which is very tasty. But no one really thinks that a fruit smoothie is a very good way to describe the original fruits, do they?

13.4.1 Doing the test in R

Doing the Welch test in R is very simple. It's exactly the same as the Student test, except that you specify `var.equal = FALSE`, or just don't bother to type in anything for the `var.equal` argument since by default it's set to `FALSE` anyway. So either of these commands will work just fine:

```
> t.test( x = grade.A, y = grade.B )
> t.test( formula = grade ~ tutor, data = harpo )
```

As with the Student test, the `formula` and `data` version produces output that is slightly different in some superficial details from the `x` and `y` version; but all of the important stuff (the numbers!) is identical. On a whim, I'll use the formula version, which means that R produces the following output:

```
Welch Two Sample t-test

data: grade by tutor
t = 2.0342, df = 23.025, p-value = 0.05361
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-0.09249349 11.04804904
sample estimates:
mean in group Anastasia mean in group Bernadette
74.53333 69.05556
```

The interpretation of this output should be fairly obvious at this point. You read the output in the same way that you would for the Student test. The most important information is all in the line that contains the *t*-statistic itself. The degrees of freedom isn't a whole number anymore, but that's to be expected given how ugly the formula for *df* was. So that's all pretty easy. Except, except... our result isn't significant anymore. When we ran the Student test, we did get a significant effect; but the Welch test on the same data set is not ($t(23.03) = 2.03, p = .054$).

What does this mean? Should we panic? Is the sky burning? Probably not. The fact that one test is significant and the other isn't doesn't itself mean very much, since I actually rigged the data so that this would happen. As a general rule, it's not a good idea to go out of your way to try to interpret or explain the difference between a *p*-value of .049 and a *p*-value of .051. If this sort of thing happens in real life, the *difference* in these *p*-values is almost certainly due to chance. What does matter is that you take a little bit of care in thinking about what test you use. The Student test and the Welch test have different strengths and weaknesses. If the two populations really do have equal variances, then the Student test is slightly more powerful (lower Type II error rate) than the Welch test. However, if they *don't* have the same variances, then the assumptions of the Student test are violated and you may not be able to trust it: you might end up with a higher Type I error rate. So it's a trade off. However, in real life, I tend to prefer the Welch test; because almost no-one *actually* believes that the population variances are identical.

13.4.2 Assumptions of the test

The assumptions of the Welch test are very similar to those made by the Student *t*-test (see Section 13.3.10), except that the Welch test does not assume homogeneity of variance. This leaves only the assumption of normality, and the assumption of independence. The specifics of these assumptions are

the same for the Welch test as for the Student test.

13.5

The paired-samples *t*-test

Regardless of whether we're talking about the Student test or the Welch test, an independent samples *t*-test is intended to be used in a situation where you have two samples that are, well, independent of one another. This situation arises naturally when participants are assigned randomly to one of two experimental conditions, but it provides a very poor approximation to other sorts of research designs. In particular, a repeated measures design – in which each participant is measured (with respect to the same outcome variable) in both experimental conditions – is not suited for analysis using independent samples *t*-tests. For example, we might be interested in whether listening to music reduces people's working memory capacity. To that end, we could measure each person's working memory capacity in two conditions: with music, and without music. In an experimental design such as this one,¹⁵ each participant appears in *both* groups. This requires us to approach the problem in a different way; by using the **paired samples *t*-test**.

13.5.1 The data

The data set that we'll use this time comes from Dr Chico's class.¹⁶ In her class, students take two major tests, one early in the semester and one later in the semester. To hear her tell it, she runs a very hard class, one that most students find very challenging; but she argues that by setting hard assessments, students are encouraged to work harder. Her theory is that the first test is a bit of a “wake up call” for students: when they realise how hard her class really is, they'll work harder for the second test and get a better mark. Is she right? To test this, let's have a look at the `chico.Rdata` file:

```
> load("chico.Rdata")
> who(TRUE)
-- Name --      -- Class --      -- Size --
chico          data.frame     20 x 3
$id            factor        20
$grade_test1   numeric       20
$grade_test2   numeric       20
```

The data frame `chico` contains three variables: an `id` variable that identifies each student in the class, the `grade_test1` variable that records the student grade for the first test, and the `grade_test2` variable that has the grades for the second test. Here's the first six students:

```
> head(chico)
  id grade_test1 grade_test2
1 student1    42.9      44.6
2 student2    51.8      54.0
3 student3    71.7      72.3
4 student4    51.6      53.4
5 student5    63.5      63.8
6 student6    58.0      59.3
```

¹⁵This design is very similar to the one in Section 12.7 that motivated the McNemar test. This should be no surprise. Both are standard repeated measures designs involving two measurements. The only difference is that this time our outcome variable is interval scale (working memory capacity) rather than a binary, nominal scale variable (a yes-or-no question).

¹⁶At this point we have Drs Harpo, Chico and Zeppo. No prizes for guessing who Dr Groucho is.

At a glance, it does seem like the class is a hard one (most grades are between 50% and 60%), but it does look like there's an improvement from the first test to the second one. If we take a quick look at the descriptive statistics

```
> library( psych )
> describe( chico )
   var n  mean    sd median trimmed  mad  min  max range skew kurtosis    se
id*      1 20 10.50 5.92   10.5  10.50 7.41   1.0 20.0 19.0  0.00 -1.20 1.32
grade_test1  2 20 56.98 6.62   57.7  56.92 7.71 42.9 71.7 28.8  0.05  0.28 1.48
grade_test2  3 20 58.38 6.41   59.7  58.35 6.45 44.6 72.3 27.7 -0.05  0.23 1.43
```

we see that this impression seems to be supported. Across all 20 students¹⁷ the mean grade for the first test is 57%, but this rises to 58% for the second test. Although, given that the standard deviations are 6.6% and 6.4% respectively, it's starting to feel like maybe the improvement is just illusory; maybe just random variation.

13.5.2 Exploring the data

Now, notice that this data set has a **repeated measures** structure: each person in the class is “measured” at two different time points. So if we want to we could convert this “wide form” data frame into a “long form”. Again, notice that I've named the variables in such a way that we can do this really easily using the `wideToLong()` function in the `lsr` package. So let's do that:

```
> chico2 <- wideToLong( data = chico,           # original data is "chico"
+                         within = "time"       # name for the grouping variable
+ )
```

Here's what the reshaped data look like

```
> head( chico2 )
  id grade time
1 student1 44.6 test2
2 student1 42.9 test1
3 student2 54.0 test2
4 student2 51.8 test1
5 student3 72.3 test2
6 student3 71.7 test1
```

As we'll see shortly, we won't actually need the data in this format in order to do our *t*-test, but it's handy for drawing some pictures, and in any case this format is very useful for other kinds of repeated measures analyses; so we might as well get into the habit of being able to switch back and forth between wide and long forms of the data.

Okay, while we've got the data in this format, let's use the `plotmeans()` function to plot the mean grade for both tests, along with 95% confidence intervals. The command to do this is:

```
> library( gplots )
> plotmeans( formula = grade ~ time,      # plot grade by test time
+            data = chico2,                 # data frame
+            n.label = FALSE,              # don't show sample size
+            xlab = "Testing Instance", # x-axis label
```

¹⁷This is obviously a class being taught at a very small or very expensive university, or else is a postgraduate class. I've never taught an intro stats class with less than 350 students.

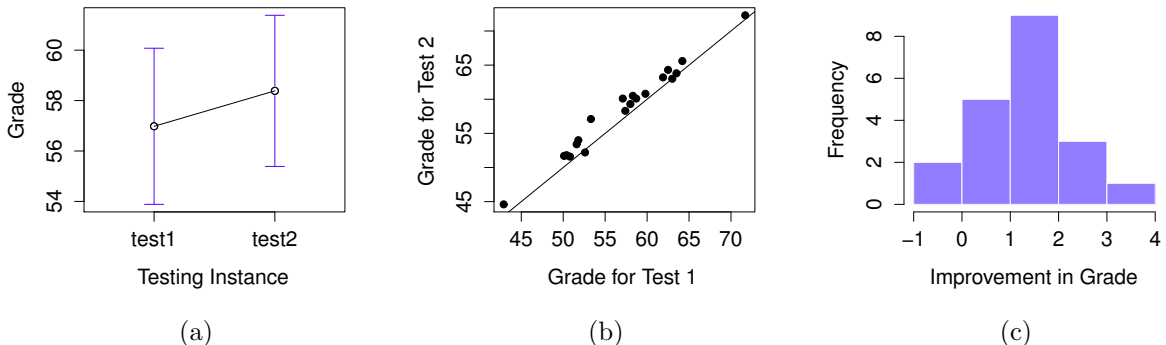


Figure 13.10: Mean grade for test 1 and test 2, with associated 95% confidence intervals (panel a). Scatterplot showing the individual grades for test 1 and test 2 (panel b). A histogram showing the distribution of improvement scores across students (panel c).

```
+           ylab = "Grade"          # y-axis label
+ )
```

and the resulting plot is shown as Figure 13.10a. If we were to rely on this plot alone, we'd come to the same conclusion that we got from looking at the descriptive statistics that the `describe()` function spat out: looking at how wide those confidence intervals are, we'd be tempted to think that the apparent improvement in student performance is pure chance.

Nevertheless, this impression is wrong. To see why, let's draw a scatterplot of the grades for test 1 against the grades for test 2. We can do this using the `plot()` command in a straightforward manner, but I'll also add a line to the plot using the `abline()` function corresponding to the line $y = x$. Here's the commands:

```
> # first draw the basic scatterplot...
> plot( x = chico$grade_test1,      # test 1 on the x-axis
+       y = chico$grade_test2,      # test 2 on the y-axis
+       xlab = "Grade for Test 1", # label for the x-axis
+       ylab = "Grade for Test 2", # label for the y-axis
+       pch = 19                  # solid dots as markers
+ )
> # now add the line...
> abline( a = 0, # line has an intercept at 0
+         b = 1  # and a slope of 1
+ )
```

The resulting plot is shown in Figure 13.10b. In this plot, each dot corresponds to the two grades for a given student: if their grade for test 1 (x co-ordinate) equals their grade for test 2 (y co-ordinate), then the dot falls on the line. Points falling above the line are the students that performed better on the second test. Critically, most of the students *do* seem to have improved their grade, if only by a small amount. To formalise this, let's create a new variable for the `improvement` that each student makes. The easiest way to do this would be to do this:

```
> improvement <- with(chico, grade_test2 - grade_test1)
```

but what this would do is create `improvement` as a variable in the global workspace. And today, what I really would prefer to do is add this to a data frame. So instead I'll use the `within()` function:

```
> chico3 <- within( chico, improvement <- grade_test2 - grade_test1)
> head( chico3 )
  id grade_test1 grade_test2 improvement
1 student1      42.9      44.6       1.7
2 student2      51.8      54.0       2.2
3 student3      71.7      72.3       0.6
4 student4      51.6      53.4       1.8
5 student5      63.5      63.8       0.3
6 student6      58.0      59.3       1.3
```

So as you can see, the `chico3` data frame now has the `improvement` variable added to it. It's not a big deal which way you do it, but I find it's nice to store new variables within a data frame whenever I can: it makes it easier to relate the new variable to the original data. In any case, now that we've created and stored this `improvement` variable, let's draw a histogram. The command I used is,

```
> hist( x = chico3$improvement,           # data to plot
+       xlab = "Improvement in Grade", # x-axis label
+       main = "",                  # no title
+       border = "white",          # white borders on the bars
+       col = rgb(.5,.5,1)         # light blue fill colour
+ )
```

and the histogram itself is Figure 13.10c. When we look at the distribution of the `improvement` scores, it's very clear that there *is* a real improvement here. The vast majority of the students scored higher on the test 2 than on test 1, reflected in the fact that almost the entire histogram is above zero. And if we now look at the confidence interval for the mean of this new variable,

```
> ciMean( x = chico3$improvement )
  2.5%    97.5%
0.9508686 1.8591314
```

we see that it is 95% certain that the true (population-wide) average improvement would lie between 0.95% and 1.86%. So you can see, qualitatively, what's going on: there is a real “within student” improvement (everyone improves by about 1%), but it is very small when set against the quite large “between student” differences (student grades vary by about 20% or so).

13.5.3 What is the paired samples t -test?

In light of the previous exploration, let's think about how to construct an appropriate t test. One possibility would be to try to run an independent samples t -test using `grade_test1` and `grade_test2` as the variables of interest. However, this is clearly the wrong thing to do: the independent samples t -test assumes that there is no particular relationship between the two samples. Yet clearly that's not true in this case, because of the repeated measures structure to the data. To use the language that I introduced in the last section, if we were to try to do an independent samples t -test, we would be conflating the `within subject` differences (which is what we're interested in testing) with the `between subject` variability (which we are not).

The solution to the problem is obvious, I hope, since we already did all the hard work in the previous section. Instead of running an independent samples t -test on `grade_test1` and `grade_test2`, we run a

one-sample t -test on the within-subject difference variable, `improvement`. To formalise this slightly, if X_{i1} is the score that the i -th participant obtained on the first variable, and X_{i2} is the score that the same person obtained on the second one, then the difference score is:

$$D_i = X_{i1} - X_{i2}$$

Notice that the difference scores is *variable 1 minus variable 2* and not the other way around, so if we want improvement to correspond to a positive valued difference, we actually want “test 2” to be our “variable 1”. Equally, we would say that $\mu_D = \mu_1 - \mu_2$ is the population mean for this difference variable. So, to convert this to a hypothesis test, our null hypothesis is that this mean difference is zero; the alternative hypothesis is that it is not:

$$\begin{aligned} H_0 : \mu_D &= 0 \\ H_1 : \mu_D &\neq 0 \end{aligned}$$

(this is assuming we’re talking about a two-sided test here). This is more or less identical to the way we described the hypotheses for the one-sample t -test: the only difference is that the specific value that the null hypothesis predicts is 0. And so our t -statistic is defined in more or less the same way too. If we let \bar{D} denote the mean of the difference scores, then

$$t = \frac{\bar{D}}{\text{SE}(\bar{D})}$$

which is

$$t = \frac{\bar{D}}{\hat{\sigma}_D / \sqrt{N}}$$

where $\hat{\sigma}_D$ is the standard deviation of the difference scores. Since this is just an ordinary, one-sample t -test, with nothing special about it, the degrees of freedom are still $N - 1$. And that’s it: the paired samples t -test really isn’t a new test at all: it’s a one-sample t -test, but applied to the difference between two variables. It’s actually very simple; the only reason it merits a discussion as long as the one we’ve just gone through is that you need to be able to recognise *when* a paired samples test is appropriate, and to understand *why* it’s better than an independent samples t test.

13.5.4 Doing the test in R

So how do we do this test in R? Obviously, it would be fairly easy to calculate the difference scores ourselves: that’s exactly what we did when creating the `improvement` variable in the first place. However, the `t.test()` function provides us with the `paired` argument, which can be used to indicate that it’s a paired test. The key arguments are thus:

- `x`. The first variable.
- `y`. The second variable
- `paired`. Set `paired = TRUE`

As usual, we can use `conf.level` to adjust the size of the confidence intervals it reports, and `alternative` to specify whether the test is one-sided or two-sided. In fact, when it comes to testing Dr Chico’s hypothesis, we genuinely do have a one-sided hypothesis: her prediction is that students *improve* their grade. Thus our null and alternative hypotheses are the one-sided versions:

$$\begin{aligned} H_0 : \mu_D &\leq 0 \\ H_1 : \mu_D &> 0 \end{aligned}$$

And thus to run my test, the command is

```

> t.test( x = chico$grade_test2,      # variable 1 is the "test2" scores
+        y = chico$grade_test1,      # variable 2 is the "test1" scores
+        paired = TRUE,             # paired test
+        alternative = "greater"  # one sided alternative: variable 1 > variable 2
+ )

```

and the output is

```

Paired t-test

data: chico$grade_test2 and chico$grade_test1
t = 6.4754, df = 19, p-value = 1.66e-06
alternative hypothesis: true difference in means is greater than 0
95 percent confidence interval:
 1.029823      Inf
sample estimates:
mean of the differences
               1.405

```

This output is very similar to the ones we've seen previously, but there's two new things, so again let's look at it closely. Firstly, let's start by noting that the important line (the one with the t statistic and the p value) is hasn't changed, so it is clear that there is a significant effect ($t(19) = 6.48, p < .001$). However, at the bottom, notice that the `sample estimates` that it reports aren't the same as before: specifically, it reports the mean of the difference scores. This is a consequence of the fact that we've done a paired samples t -test, rather than an independent samples test.

The second thing that is new is the confidence interval, which now ranges from 1.03 to ∞ (i.e., infinity). This change has nothing to do with the fact that we did a paired samples test: it's because we're doing a one-sided test. Because we have one-sided hypotheses, R also reports a one-sided confidence interval. That is, what it reports is a single number (in this case 1.03); where we can feel 95% confident that $\mu_D > 1.03$.

13.5.5 A flaw in the design of the `t.test()` function

As far as the actual statistics go, we're now done with t -tests. However, there's one last thing I need to point out about the `t.test()` function itself. There is a tiny bug in the way that the `t.test()` function is written. This bug won't affect the results of any of your actual t -tests, so there's nothing to worry about statistically. It's just that `t.test()` violates one of the usual rules about arguments and functions. Remember how, way back in Section 3.5.1 I explained that R doesn't care about the order in which you specify the arguments, as long as you use the argument names? The `t.test()` function violates that rule, but only if you are using the `formula` argument to specify your t -test. The `formula` argument must come first, regardless of whether you use the name. As far as I can tell, this is a genuine mistake in the design

of the function.¹⁸

13.6 _____

Effect size

The most commonly used measure of effect size for a t -test is **Cohen's d** (Cohen, 1988). It's a very simple measure in principle, with quite a few wrinkles when you start digging into the details. Cohen himself defined it primarily in the context of an independent samples t -test, specifically the Student test. In that context, a natural way of defining the effect size is to divide the difference between the means by an estimate of the standard deviation. In other words, we're looking to calculate *something* along the lines of this:

$$\frac{(\text{mean 1}) - (\text{mean 2})}{\text{std dev}}$$

You'd think that this would be an unambiguous measure, but it's not; largely because Cohen wasn't too specific on what he thought should be used as the measure of the standard deviation (in his defence, he was trying to make a broader point in his book, not nitpick about tiny details). As discussed by McGrath and Meyer (2006), there are several different version in common usage, and each author tends to adopt slightly different notation. For the sake of simplicity (as opposed to accuracy) I'll use d to refer to any statistic that you calculate from the sample, and use δ to refer to a theoretical population effect. Obviously, that does mean that there are several different things all called d . However, the `cohensD()` function in the `lsr` package uses the `method` argument to distinguish between them, so that's what I'll do in the text.

13.6.1 Cohen's d from one sample

The simplest situation to consider is the one corresponding to a one-sample t -test. In this case, the one sample mean \bar{X} and one (hypothesised) population mean μ_0 to compare it to. Not only that, there's really only one sensible way to estimate the population standard deviation: we just use our usual estimate $\hat{\sigma}$. Therefore, we end up with the following as the only way to calculate d ,

$$d = \frac{\bar{X} - \mu_0}{\hat{\sigma}}$$

To calculate it in R, you can use the `cohensD()` function from the `lsr` package. When writing this function, I've made some attempt to make it work in a similar way to `t.test()`. As a consequence, `cohensD()` can calculate your effect size regardless of which type of t -test you performed. If what you want is a measure of Cohen's d to accompany a one-sample t -test, there's only two arguments that you need to care about. These are:

- `x`. A numeric vector containing the sample data.
- `mu`. The mean against which the mean of `x` is compared (default value is `mu = 0`).

We don't need to specify what `method` to use, because there's only one version of d that makes sense in this context. So, in order to compute an effect size for the data from Dr Zeppo's class (Section 13.2), we'd type something like this:

¹⁸Computer geeks only: the reason for this is that method dispatch in the S3 object oriented system is a little ad hoc, and is based either on the first argument or by the name of an argument specified in the `UseMethod()` function. The problem with `t.test()` is that method dispatch is based on `x` and not `formula`. If `formula` is the first argument and there's no `x` value then everything is fine, but if there's a `formula` argument somewhere else then R passes everything to the default method, which doesn't take `formula` inputs. Sigh.

```

> cohensD( x = grades,      # data are stored in the grades vector
+           mu = 67.5       # compare students to a mean of 67.5
+ )
[1] 0.5041691

```

and, just so that you can see that there's nothing fancy going on, the command below shows you how to calculate it if there weren't no fancypants `cohensD()` function available:

```

> ( mean(grades) - 67.5 ) / sd(grades)
[1] 0.5041691

```

Yep, same number. Overall, then, the psychology students in Dr Zeppo's class are achieving grades (mean = 72.3%) that are about .5 standard deviations higher than the level that you'd expect (67.5%) if they were performing at the same level as other students. Judged against Cohen's rough guide, this is a moderate effect size.

13.6.2 Cohen's d from a Student t test

The majority of discussions of Cohen's d focus on a situation that is analogous to Student's independent samples t test, and it's in this context that the story becomes messier, since there are several different versions of d that you might want to use in this situation, and you can use the `method` argument to the `cohensD()` function to pick the one you want. To understand why there are multiple versions of d , it helps to take the time to write down a formula that corresponds to the true population effect size δ . It's pretty straightforward,

$$\delta = \frac{\mu_1 - \mu_2}{\sigma}$$

where, as usual, μ_1 and μ_2 are the population means corresponding to group 1 and group 2 respectively, and σ is the standard deviation (the same for both populations). The obvious way to estimate δ is to do exactly the same thing that we did in the t -test itself: use the sample means as the top line, and a pooled standard deviation estimate for the bottom line:

$$d = \frac{\bar{X}_1 - \bar{X}_2}{\hat{\sigma}_p}$$

where $\hat{\sigma}_p$ is the exact same pooled standard deviation measure that appears in the t -test. This is the most commonly used version of Cohen's d when applied to the outcome of a Student t -test ,and is sometimes referred to as Hedges' g statistic (Hedges, 1981). It corresponds to `method = "pooled"` in the `cohensD()` function, and it's the default.

However, there are other possibilities, which I'll briefly describe. Firstly, you may have reason to want to use only one of the two groups as the basis for calculating the standard deviation. This approach (often called Glass' Δ) only makes most sense when you have good reason to treat one of the two groups as a purer reflection of "natural variation" than the other. This can happen if, for instance, one of the two groups is a control group. If that's what you want, then use `method = "x.sd"` or `method = "y.sd"` when using `cohensD()`. Secondly, recall that in the usual calculation of the pooled standard deviation we divide by $N - 2$ to correct for the bias in the sample variance; in one version of Cohen's d this correction is omitted. Instead, we divide by N . This version (`method = "raw"`) makes sense primarily when you're trying to calculate the effect size in the sample; rather than estimating an effect size in the population. Finally, there is a version based on Hedges and Olkin (1985), who point out there is a small bias in the usual (pooled) estimation for Cohen's d . Thus they introduce a small correction (`method = "corrected"`), by multiplying the usual value of d by $(N - 3)/(N - 2.25)$.

In any case, ignoring all those variations that you could make use of if you wanted, let's have a look at how to calculate the default version. In particular, suppose we look at the data from the `harpo2.Rdata` file, in which the data from Anastasia's class are stored in `grade.A` and the data from Bernadette's class are stored in the `grade.B` variable. The relevant arguments in this context are

- `x`. Numeric vector with data from group 1.
- `y`. Numeric vector with data from group 2.
- `method`. A character string indicating which version you want to calculate. The values that make sense for the Student test are `"pooled"` (the default), `"x.sd"`, `"y.sd"`, `"raw"` and `"corrected"`. The differences between these are as described above.

and so we can calculate Cohen's d like this:

```
> cohensD( x = grade.A,      # data from Anastasia's tutorial
+           y = grade.B,      # data from Bernadette's tutorial
+           method = "pooled" # which version to calculate?
+ )
[1] 0.7395614
```

although, we really didn't need to include the `method = "pooled"` part, since this is the default method. However, that's not the only way to do it. Just like we saw with the Student t -test itself, you're much more likely to have the data stored in a data frame like the `harpo` one that we have saved in the `harpo.Rdata` file. To make our lives easier, we can specify `formula` and `data` instead of `x` and `y`.

- `formula`. A formula of the format `outcome ~ group`. That is, on the left hand side you type the name of the outcome variable, and on the right hand side you type the name of the grouping variable.
- `data`. The data frame where the variables are stored. If this argument isn't specified, R will look in the workspace to see if the variables are there.

That means that the following is an equivalent way to do the calculations

```
> cohensD( formula = grade ~ tutor,  # outcome ~ group
+           data = harpo,          # data frame
+           method = "pooled"       # which version to calculate?
+ )
[1] 0.7395614
```

13.6.3 Cohen's d from a Welch test

Suppose the situation you're in is more like the Welch test: you still have two independent samples, but you no longer believe that the corresponding populations have equal variances. When this happens, we have to redefine what we mean by the population effect size. I'll refer to this new measure as δ' , so as to keep it distinct from the measure δ which we defined previously. What Cohen (1988) suggests is that we could define our new population effect size by averaging the two population variances. What this means is that we get:

$$\delta' = \frac{\mu_1 - \mu_2}{\sigma'}$$

where

$$\sigma' = \sqrt{\frac{\sigma_1^2 + \sigma_2^2}{2}}$$

This seems quite reasonable, but notice that none of the measures that we've discussed so far are attempting to estimate this new quantity. It might just be my own ignorance of the topic, but I'm only aware of one version of Cohen's d that actually estimates the unequal-variance effect size δ' rather than the equal-variance effect size δ . All we do to calculate d for this version (`method = "unequal"`) is substitute the sample means \bar{X}_1 and \bar{X}_2 and the corrected sample standard deviations $\hat{\sigma}_1$ and $\hat{\sigma}_2$ into the equation for δ' . This gives us the following equation for d ,

$$d = \frac{\bar{X}_1 - \bar{X}_2}{\sqrt{\frac{\hat{\sigma}_1^2 + \hat{\sigma}_2^2}{2}}}$$

as our estimate of the effect size. There's nothing particularly difficult about calculating this version in R, since all we have to do is change the `method` argument:

```
> cohensD( formula = grade ~ tutor,
+           data = harpo,
+           method = "unequal"
+ )
[1] 0.7244995
```

13.6.4 Cohen's d from a paired-samples test

Finally, what should we do for a paired samples t -test? In this case, the answer depends on what it is you're trying to do. If you want to measure your effect sizes relative to the distribution of difference scores, the measure of d that you calculate is just (`method = "paired"`)

$$d = \frac{\bar{D}}{\hat{\sigma}_D}$$

where $\hat{\sigma}_D$ is the estimate of the standard deviation of the differences. The calculation here is pretty straightforward

```
> cohensD( x = chico$grade_test2,
+           y = chico$grade_test1,
+           method = "paired"
+ )
[1] 1.447952
```

The only wrinkle is figuring out whether this is the measure you want or not. To the extent that you care about the practical consequences of your research, you often want to measure the effect size relative to the *original* variables, not the *difference* scores (e.g., the 1% improvement in Dr Chico's class is pretty small when measured against the amount of between-student variation in grades), in which case you use the same versions of Cohen's d that you would use for a Student or Welch test. For instance, when we do that for Dr Chico's class,

```
> cohensD( x = chico$grade_test2,
+           y = chico$grade_test1,
+           method = "pooled"
+ )
[1] 0.2157646
```

what we see is that the overall effect size is quite small, when assessed on the scale of the original variables.

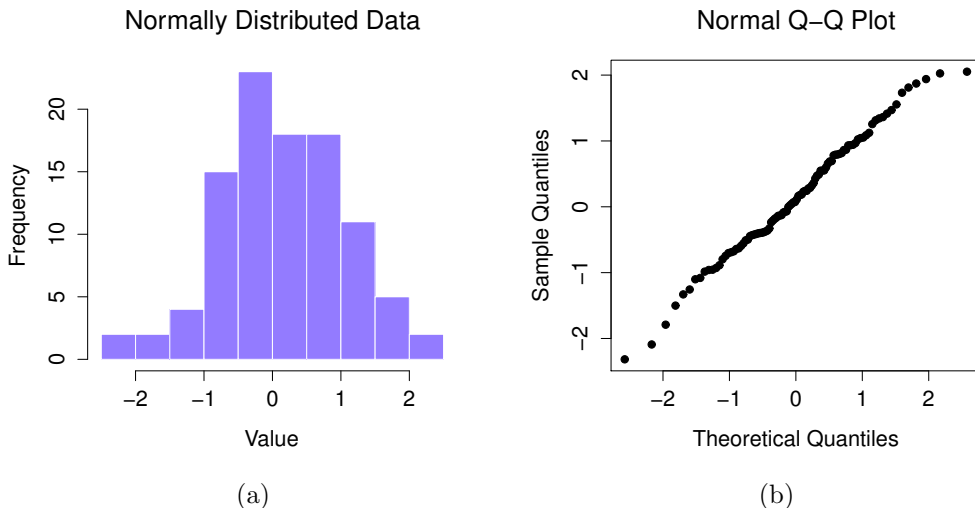


Figure 13.11: Histogram (panel a) and normal QQ plot (panel b) of `normal.data`, a normally distributed sample with 100 observations. The Shapiro-Wilk statistic associated with these data is $W = .99$, indicating that no significant departures from normality were detected ($p = .73$).

13.7

Checking the normality of a sample

All of the tests that we have discussed so far in this chapter have assumed that the data are normally distributed. This assumption is often quite reasonable, because the central limit theorem (Section 10.4) does tend to ensure that many real world quantities are normally distributed: any time that you suspect that your variable is *actually* an average of lots of different things, there's a pretty good chance that it will be normally distributed; or at least close enough to normal that you can get away with using t -tests. However, life doesn't come with guarantees; and besides, there are lots of ways in which you can end up with variables that are highly non-normal. For example, any time you think that your variable is actually the minimum of lots of different things, there's a very good chance it will end up quite skewed. In psychology, response time (RT) data is a good example of this. If you suppose that there are lots of things that could trigger a response from a human participant, then the actual response will occur the first time one of these trigger events occurs.¹⁹ This means that RT data are systematically non-normal. Okay, so if normality is assumed by all the tests, and is mostly but not always satisfied (at least approximately) by real world data, how can we check the normality of a sample? In this section I discuss two methods: QQ plots, and the Shapiro-Wilk test.

13.7.1 QQ plots

One way to check whether a sample violates the normality assumption is to draw a “**quantile-quantile**” plot (QQ plot). This allows you to visually check whether you’re seeing any systematic violations. In a

¹⁹This is a massive oversimplification.

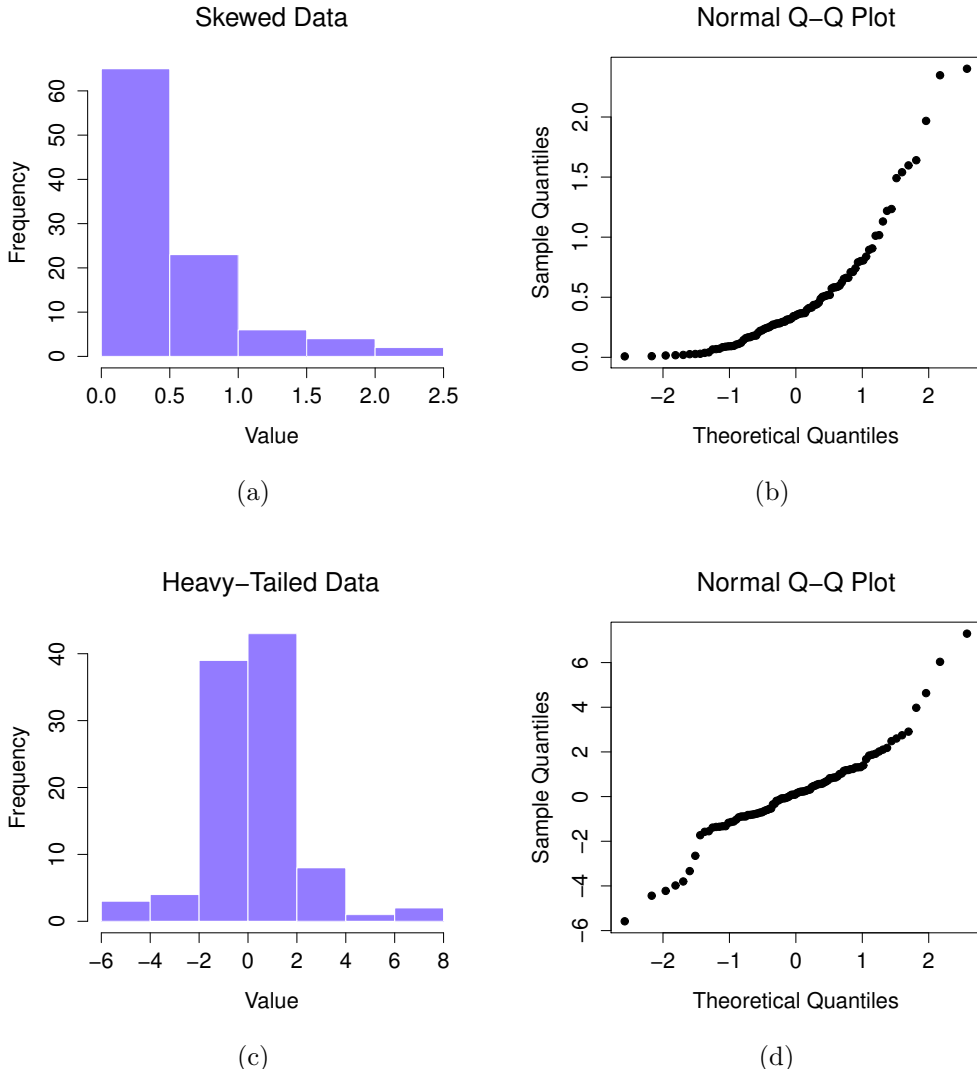


Figure 13.12: In the top row, a histogram (panel a) and normal QQ plot (panel b) of the 100 observations in a `skewed.data` set. The skewness of the data here is 1.94, and is reflected in a QQ plot that curves upwards. As a consequence, the Shapiro-Wilk statistic is $W = .80$, reflecting a significant departure from normality ($p < .001$). The bottom row shows the same plots for a heavy tailed data set, again consisting of 100 observations. In this case, the heavy tails in the data produce a high kurtosis (2.80), and cause the QQ plot to flatten in the middle, and curve away sharply on either side. The resulting Shapiro-Wilk statistic is $W = .93$, again reflecting significant non-normality ($p < .001$).

QQ plot, each observation is plotted as a single dot. The x co-ordinate is the theoretical quantile that the observation should fall in, if the data were normally distributed (with mean and variance estimated from the sample) and on the y co-ordinate is the actual quantile of the data within the sample. If the data are normal, the dots should form a straight line. For instance, lets see what happens if we generate data by sampling from a normal distribution, and then drawing a QQ plot using the R function `qqnorm()`. The `qqnorm()` function has a few arguments, but the only one we really need to care about here is `y`, a vector specifying the data whose normality we're interested in checking. Here's the R commands:

```
> normal.data <- rnorm( n = 100 ) # generate N = 100 normally distributed numbers
> hist( x = normal.data )          # draw a histogram of these numbers
> qqnorm( y = normal.data )       # draw the QQ plot
```

And the results are shown in Figure 13.11. As you can see, these data form a pretty straight line; which is no surprise given that we sampled them from a normal distribution! In contrast, have a look at the two data sets shown in Figure 13.12. The top panels show the histogram and a QQ plot for a data set that is highly skewed: the QQ plot curves upwards. The lower panels show the same plots for a heavy tailed (i.e., high kurtosis) data set: in this case, the QQ plot flattens in the middle and curves sharply at either end.

13.7.2 Shapiro-Wilk tests

Although QQ plots provide a nice way to informally check the normality of your data, sometimes you'll want to do something a bit more formal. And when that moment comes, the **Shapiro-Wilk test** (Shapiro & Wilk, 1965) is probably what you're looking for.²⁰ As you'd expect, the null hypothesis being tested is that a set of N observations is normally distributed. The test statistic that it calculates is conventionally denoted as W , and it's calculated as follows. First, we sort the observations in order of increasing size, and let X_1 be the smallest value in the sample, X_2 be the second smallest and so on. Then the value of W is given by

$$W = \frac{\left(\sum_{i=1}^N a_i X_i\right)^2}{\sum_{i=1}^N (X_i - \bar{X})^2}$$

where \bar{X} is the mean of the observations, and the a_i values are ... mumble, mumble ... something complicated that is a bit beyond the scope of an introductory text.

Because it's a little hard to explain the maths behind the W statistic, a better idea is to give a broad brush description of how it behaves. Unlike most of the test statistics that we'll encounter in this book, it's actually *small* values of W that indicated departure from normality. The W statistic has a maximum value of 1, which arises when the data look "perfectly normal". The smaller the value of W , the less normal the data are. However, the sampling distribution for W – which is not one of the standard ones that I discussed in Chapter 9 and is in fact a complete pain in the arse to work with – does depend on the sample size N . To give you a feel for what these sampling distributions look like, I've plotted three of them in Figure 13.13. Notice that, as the sample size starts to get large, the sampling distribution becomes very tightly clumped up near $W = 1$, and as a consequence, for larger samples W doesn't have to be very much smaller than 1 in order for the test to be significant.

To run the test in R, we use the `shapiro.test()` function. It has only a single argument `x`, which is a numeric vector containing the data whose normality needs to be tested. For example, when we apply this function to our `normal.data`, we get the following:

²⁰Either that, or the Kolmogorov-Smirnov test, which is probably more traditional than the Shapiro-Wilk, though most things I've read seem to suggest Shapiro-Wilk is the better test of normality; although Kolomogorov-Smirnov is a general purpose test of distributional equivalence, so it can be adapted to handle other kinds of distribution tests; in R it's implemented via the `ks.test()` function.

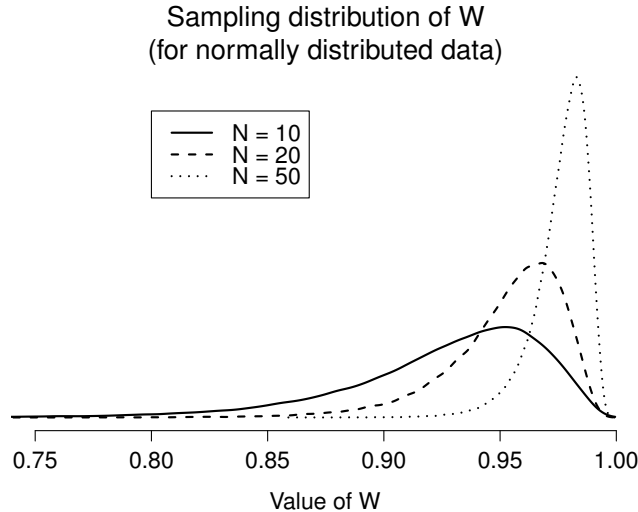


Figure 13.13: Sampling distribution of the Shapiro-Wilk W statistic, under the null hypothesis that the data are normally distributed, for samples of size 10, 20 and 50. Note that *small* values of W indicate departure from normality.

```
> shapiro.test( x = normal.data )

Shapiro-Wilk normality test

data: normal.data
W = 0.9903, p-value = 0.6862
```

So, not surprisingly, we have no evidence that these data depart from normality. When reporting the results for a Shapiro-Wilk test, you should (as usual) make sure to include the test statistic W and the p value, though given that the sampling distribution depends so heavily on N it would probably be a politeness to include N as well.

13.8

Testing non-normal data with Wilcoxon tests

Okay, suppose your data turn out to be pretty substantially non-normal, but you still want to run something like a t -test? This situation occurs a lot in real life: for the AFL winning margins data, for instance, the Shapiro-Wilk test made it very clear that the normality assumption is violated. This is the situation where you want to use Wilcoxon tests.

Like the t -test, the Wilcoxon test comes in two forms, one-sample and two-sample, and they're used in more or less the exact same situations as the corresponding t -tests. Unlike the t -test, the Wilcoxon test doesn't assume normality, which is nice. In fact, they don't make any assumptions about what kind of distribution is involved: in statistical jargon, this makes them **nonparametric tests**. While avoiding the normality assumption is nice, there's a drawback: the Wilcoxon test is usually less powerful than the

t-test (i.e., higher Type II error rate). I won't discuss the Wilcoxon tests in as much detail as the *t*-tests, but I'll give you a brief overview.

13.8.1 Two sample Wilcoxon test

I'll start by describing the **two sample Wilcoxon test** (also known as the Mann-Whitney test), since it's actually simpler than the one sample version. Suppose we're looking at the scores of 10 people on some test. Since my imagination has now failed me completely, let's pretend it's a "test of awesomeness", and there are two groups of people, "A" and "B". I'm curious to know which group is more awesome. The data are included in the file `awesome.Rdata`, and like many of the data sets I've been using, it contains only a single data frame, in this case called `awesome`. Here's the data:

```
> load("awesome.Rdata")
> print( awesome )
  scores group
  1     6.4    A
  2    10.7    A
  3   11.9    A
  4    7.3    A
  5   10.0    A
  6   14.5    B
  7   10.4    B
  8   12.9    B
  9   11.7    B
 10   13.0    B
```

As long as there are no ties (i.e., people with the exact same awesomeness score), then the test that we want to do is surprisingly simple. All we have to do is construct a table that compares every observation in group *A* against every observation in group *B*. Whenever the group *A* datum is larger, we place a check mark in the table:

		group B				
		14.5	10.4	12.4	11.7	13.0
group A	6.4
	10.7	.	✓	.	.	.
	11.9	.	✓	.	✓	.
	7.3
	10.0

We then count up the number of checkmarks. This is our test statistic, W .²¹ The actual sampling distribution for W is somewhat complicated, and I'll skip the details. For our purposes, it's sufficient to note that the interpretation of W is qualitatively the same as the interpretation of t or z . That is, if we want a two-sided test, then we reject the null hypothesis when W is very large or very small; but if we have a directional (i.e., one-sided) hypothesis, then we only use one or the other.

The structure of the `wilcox.test()` function should feel very familiar to you by now. When you have your data organised in terms of an outcome variable and a grouping variable, then you use the `formula` and `data` arguments, so your command looks like this:

²¹Actually, there are two different versions of the test statistic; they differ from each other by a constant value. The version that I've described is the one that R calculates.

```

> wilcox.test( formula = scores ~ group, data = awesome)

Wilcoxon rank sum test

data: scores by group
W = 3, p-value = 0.05556
alternative hypothesis: true location shift is not equal to 0

```

Just like we saw with the `t.test()` function, there is an `alternative` argument that you can use to switch between two-sided tests and one-sided tests, plus a few other arguments that we don't need to worry too much about at an introductory level. Similarly, the `wilcox.test()` function allows you to use the `x` and `y` arguments when you have your data stored separately for each group. For instance, suppose we use the data from the `awesome2.Rdata` file:

```

> load( "awesome2.Rdata" )
> score.A
[1] 6.4 10.7 11.9 7.3 10.0
> score.B
[1] 14.5 10.4 12.9 11.7 13.0

```

When your data are organised like this, then you would use a command like this:

```
> wilcox.test( x = score.A, y = score.B )
```

The output that R produces is pretty much the same as last time.

13.8.2 One sample Wilcoxon test

What about the **one sample Wilcoxon test** (or equivalently, the paired samples Wilcoxon test)? Suppose I'm interested in finding out whether taking a statistics class has any effect on the happiness of students. Here's my data:

```

> load( "happy.Rdata" )
> print( happiness )
   before after change
1      30     6    -24
2      43    29    -14
3      21    11    -10
4      24    31      7
5      23    17      -6
6      40     2    -38
7      29    31      2
8      56    21    -35
9      38     8    -30
10     16    21      5

```

What I've measured here is the happiness of each student `before` taking the class and `after` taking the class; the `change` score is the difference between the two. Just like we saw with the *t*-test, there's no fundamental difference between doing a paired-samples test using `before` and `after`, versus doing a one-sample test using the `change` scores. As before, the simplest way to think about the test is to construct a tabulation. The way to do it this time is to take those change scores that are positive valued, and tabulate them against all the complete sample. What you end up with is a table that looks like this:

		all differences									
		-24	-14	-10	7	-6	-38	2	-35	-30	5
positive differences	7	.	.	.	✓	✓	.	✓	.	.	✓
	2	✓	.	.	.
	5	✓	.	.	✓

Counting up the tick marks this time, we get a test statistic of $V = 7$. As before, if our test is two sided, then we reject the null hypothesis when V is very large or very small. As far as running it in R goes, it's pretty much what you'd expect. For the one-sample version, the command you would use is

```
> wilcox.test( x = happiness$change,
+               mu = 0
+ )
```

and the output that you get is

```
Wilcoxon signed rank test

data:  happiness$change
V = 7, p-value = 0.03711
alternative hypothesis: true location is not equal to 0
```

As this shows, we have a significant effect. Evidently, taking a statistics class does have an effect on your happiness. Switching to a paired samples version of the test won't give us different answers, of course; but here's the command to do it:

```
> wilcox.test( x = happiness$after,
+               y = happiness$before,
+               paired = TRUE
+ )
```

13.9

Summary

13.9.1 Key concepts

- A one sample t -test is used to compare a single sample mean against a hypothesised value for the population mean. (Section 13.2)
- An independent samples t -test is used to compare the means of two groups, and tests the null hypothesis that they have the same mean. It comes in two forms: the Student test (Section 13.3) assumes that the groups have the same standard deviation, the Welch test (Section 13.4) does not.
- A paired samples t -test is used when you have two scores from each person, and you want to test the null hypothesis that the two scores have the same mean. It is equivalent to taking the difference between the two scores for each person, and then running a one sample t -test on the difference scores. (Section 13.5)
- Effect size calculations for the difference between means can be calculated via the Cohen's d statistic. (Section 13.6).

- You can check the normality of a sample using QQ plots and the Shapiro-Wilk test. (Section 13.7)
- If your data are non-normal, you can use Wilcoxon tests instead of *t*-tests. (Section 13.8)

14. Comparing several means (one-way ANOVA)

This chapter introduces one of the most widely used tools in statistics, known as “the analysis of variance”, which is usually referred to as ANOVA. The basic technique was developed by Sir Ronald Fisher in the early 20th century, and it is to him that we owe the rather unfortunate terminology. The term ANOVA is a little misleading, in two respects. Firstly, although the name of the technique refers to variances, ANOVA is concerned with investigating differences in means. Secondly, there are several different things out there that are all referred to as ANOVAs, some of which have only a very tenuous connection to one another. Later on in the book we’ll encounter a range of different ANOVA methods that apply in quite different situations, but for the purposes of this chapter we’ll only consider the simplest form of ANOVA, in which we have several different groups of observations, and we’re interested in finding out whether those groups differ in terms of some outcome variable of interest. This is the question that is addressed by a **one-way ANOVA**. As with all of the chapters in this book, there are quite a few different sources that I’ve relied upon, but the one stand-out text that I’ve been most heavily influenced by is [Sahai and Ageel \(2000\)](#). It’s not a good book for beginners, but it’s an excellent book for more advanced readers who are interested in understanding the mathematics behind ANOVA.

The structure of this chapter is as follows: In Section 14.1 I’ll introduce a fictitious data set that we’ll use as a running example throughout the chapter. After introducing the data, I’ll describe the mechanics of how a one-way ANOVA actually works (Section 14.2) and then focus on how you can run one in R (Section 14.3). These two sections are the core of the chapter. The remainder of the chapter discusses a range of important topics that inevitably arise when running an ANOVA, namely how to calculate effect sizes (Section 14.4), post hoc tests and corrections for multiple comparisons (Section 14.5) and the assumptions that ANOVA relies upon (Section 14.6). We’ll also talk about how to check those assumptions and some of the things you can do if the assumptions are violated (Sections 14.7 to 14.10). At the end of the chapter we’ll talk a little about the relationship between ANOVA and other statistical tools (Section 14.11).

14.1

An illustrative data set

Suppose you’ve become involved in a clinical trial in which you are testing a new antidepressant drug called *Joyzepam*. In order to construct a fair test of the drug’s effectiveness, the study involves three separate drugs to be administered. One is a placebo, and the other is an existing antidepressant / anti-anxiety drug called *Anxifree*. A collection of 18 participants with moderate to severe depression are recruited for your initial testing. Because the drugs are sometimes administered in conjunction with psychological therapy, your study includes 9 people undergoing cognitive behavioural therapy (CBT) and 9 who are not. Participants are randomly assigned (doubly blinded, of course) a treatment, such

that there are 3 CBT people and 3 no-therapy people assigned to each of the 3 drugs. A psychologist assesses the mood of each person after a 3 month run with each drug: and the overall *improvement* in each person's mood is assessed on a scale ranging from -5 to $+5$.

With that as the study design, let's now look at what we've got in the data file:

```
> load( "clinicaltrial.Rdata" ) # load data
> who( expand = TRUE )    # inspect the workspace
-- Name --   -- Class --   -- Size --
clin.trial   data.frame   18 x 3
$drug        factor       18
$therapy     factor       18
$mood.gain  numeric      18
```

So we have a single data frame called `clin.trial`, containing three variables; `drug`, `therapy` and `mood.gain`. Next, let's print the data frame to get a sense of what the data actually look like.

```
> print( clin.trial )
    drug   therapy mood.gain
1 placebo no.therapy  0.5
2 placebo no.therapy  0.3
3 placebo no.therapy  0.1
4 anxifree no.therapy 0.6
5 anxifree no.therapy  0.4
6 anxifree no.therapy  0.2
7 joyzepam no.therapy 1.4
8 joyzepam no.therapy  1.7
9 joyzepam no.therapy  1.3
10 placebo      CBT  0.6
11 placebo      CBT  0.9
12 placebo      CBT  0.3
13 anxifree      CBT  1.1
14 anxifree      CBT  0.8
15 anxifree      CBT  1.2
16 joyzepam      CBT  1.8
17 joyzepam      CBT  1.3
18 joyzepam      CBT  1.4
```

For the purposes of this chapter, what we're really interested in is the effect of `drug` on `mood.gain`. The first thing to do is calculate some descriptive statistics and draw some graphs. In Chapter 5 we discussed a variety of different functions that can be used for this purpose. For instance, we can use the `xtabs()` function to see how many people we have in each group:

```
> xtabs( ~drug, clin.trial )
drug
placebo  anxifree  joyzepam
       6         6         6
```

Similarly, we can use the `aggregate()` function to calculate means and standard deviations for the `mood.gain` variable broken down by which `drug` was administered:

```
> aggregate( mood.gain ~ drug, clin.trial, mean )
    drug mood.gain
1 placebo 0.4500000
2 anxifree 0.7166667
```

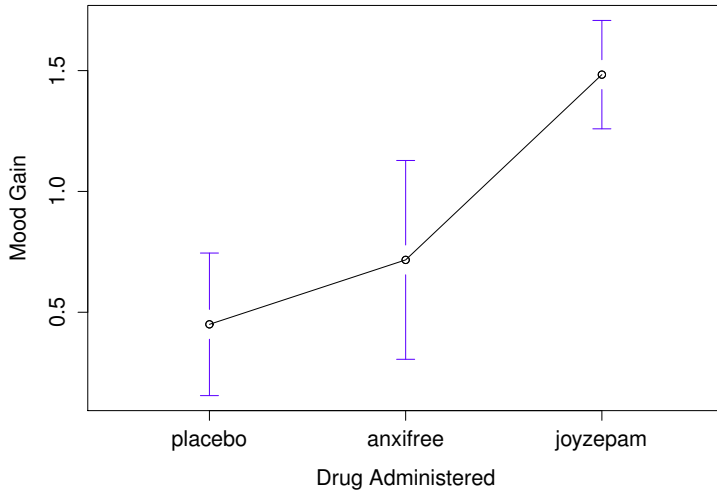


Figure 14.1: Average mood gain as a function of drug administered. Error bars depict 95% confidence intervals associated with each of the group means.

```

3 joyzepam 1.4833333

> aggregate( mood.gain ~ drug, clin.trial, sd )
   drug mood.gain
1 placebo 0.2810694
2 anxitfree 0.3920034
3 joyzepam 0.2136976

```

Finally, we can use `plotmeans()` from the `gplots` package to produce a pretty picture.

```

> library(gplots)
> plotmeans( formula = mood.gain ~ drug, # plot mood.gain by drug
+             data = clin.trial,           # the data frame
+             xlab = "Drug Administered", # x-axis label
+             ylab = "Mood Gain",        # y-axis label
+             n.label = FALSE            # don't display sample size
+ )

```

The results are shown in Figure 14.1, which plots the average mood gain for all three conditions; error bars show 95% confidence intervals. As the plot makes clear, there is a larger improvement in mood for participants in the Joyzepam group than for either the Anxitfree group or the placebo group. The Anxitfree group shows a larger mood gain than the control group, but the difference isn't as large.

The question that we want to answer is: are these difference “real”, or are they just due to chance?

14.2

How ANOVA works

In order to answer the question posed by our clinical trial data, we're going to run a one-way ANOVA. As usual, I'm going to start by showing you how to do it the hard way, building the statistical tool from the ground up and showing you how you could do it in R if you didn't have access to any of the cool built-in ANOVA functions. And, as always, I hope you'll read it carefully, try to do it the long way once or twice to make sure you really understand how ANOVA works, and then – once you've grasped the concept – never *ever* do it this way again.

The experimental design that I described in the previous section strongly suggests that we're interested in comparing the average mood change for the three different drugs. In that sense, we're talking about an analysis similar to the *t*-test (Chapter 13), but involving more than two groups. If we let μ_P denote the population mean for the mood change induced by the placebo, and let μ_A and μ_J denote the corresponding means for our two drugs, Anxifree and Joyzepam, then the (somewhat pessimistic) null hypothesis that we want to test is that all three population means are identical: that is, *neither* of the two drugs is any more effective than a placebo. Mathematically, we write this null hypothesis like this:

$$H_0 : \text{it is true that } \mu_P = \mu_A = \mu_J$$

As a consequence, our alternative hypothesis is that at least one of the three different treatments is different from the others. It's a little trickier to write this mathematically, because (as we'll discuss) there are quite a few different ways in which the null hypothesis can be false. So for now we'll just write the alternative hypothesis like this:

$$H_1 : \text{it is not true that } \mu_P = \mu_A = \mu_J$$

This null hypothesis is a lot trickier to test than any of the ones we've seen previously. How shall we do it? A sensible guess would be to "do an ANOVA", since that's the title of the chapter, but it's not particularly clear why an "analysis of *variances*" will help us learn anything useful about the *means*. In fact, this is one of the biggest conceptual difficulties that people have when first encountering ANOVA. To see how this works, I find it most helpful to start by talking about variances. In fact, what I'm going to do is start by playing some mathematical games with the formula that describes the variance. That is, we'll start out by playing around with variances, and it will turn out that this gives us a useful tool for investigating means.

14.2.1 Two formulas for the variance of Y

Firstly, let's start by introducing some notation. We'll use G to refer to the total number of groups. For our data set, there are three drugs, so there are $G = 3$ groups. Next, we'll use N to refer to the total sample size: there are a total of $N = 18$ people in our data set. Similarly, let's use N_k to denote the number of people in the k -th group. In our fake clinical trial, the sample size is $N_k = 6$ for all three groups.¹ Finally, we'll use Y to denote the outcome variable: in our case, Y refers to mood change. Specifically, we'll use Y_{ik} to refer to the mood change experienced by the i -th member of the k -th group. Similarly, we'll use \bar{Y} to be the average mood change, taken across all 18 people in the experiment, and \bar{Y}_k to refer to the average mood change experienced by the 6 people in group k .

¹When all groups have the same number of observations, the experimental design is said to be "balanced". Balance isn't such a big deal for one-way ANOVA, which is the topic of this chapter. It becomes more important when you start doing more complicated ANOVAs.

Excellent. Now that we've got our notation sorted out, we can start writing down formulas. To start with, let's recall the formula for the variance that we used in Section 5.2, way back in those kinder days when we were just doing descriptive statistics. The sample variance of Y is defined as follows:

$$\text{Var}(Y) = \frac{1}{N} \sum_{k=1}^G \sum_{i=1}^{N_k} (Y_{ik} - \bar{Y})^2$$

This formula looks pretty much identical to the formula for the variance in Section 5.2. The only difference is that this time around I've got two summations here: I'm summing over groups (i.e., values for k) and over the people within the groups (i.e., values for i). This is purely a cosmetic detail: if I'd instead used the notation Y_p to refer to the value of the outcome variable for person p in the sample, then I'd only have a single summation. The only reason that we have a double summation here is that I've classified people into groups, and then assigned numbers to people within groups.

A concrete example might be useful here. Let's consider this table, in which we have a total of $N = 5$ people sorted into $G = 2$ groups. Arbitrarily, let's say that the "cool" people are group 1, and the "uncool" people are group 2, and it turns out that we have three cool people ($N_1 = 3$) and two uncool people ($N_2 = 2$).

name	person p	group	group num. k	index in group i	grumpiness Y_{ik} or Y_p
Ann	1	cool	1	1	20
Ben	2	cool	1	2	55
Cat	3	cool	1	3	21
Dan	4	uncool	2	1	91
Egg	5	uncool	2	2	22

Notice that I've constructed two different labelling schemes here. We have a "person" variable p , so it would be perfectly sensible to refer to Y_p as the grumpiness of the p -th person in the sample. For instance, the table shows that Dan is the four so we'd say $p = 4$. So, when talking about the grumpiness Y of this "Dan" person, whoever he might be, we could refer to his grumpiness by saying that $Y_p = 91$, for person $p = 4$ that is. However, that's not the only way we could refer to Dan. As an alternative we could note that Dan belongs to the "uncool" group ($k = 2$), and is in fact the first person listed in the uncool group ($i = 1$). So it's equally valid to refer to Dan's grumpiness by saying that $Y_{ik} = 91$, where $k = 2$ and $i = 1$. In other words, each person p corresponds to a unique ik combination, and so the formula that I gave above is actually identical to our original formula for the variance, which would be

$$\text{Var}(Y) = \frac{1}{N} \sum_{p=1}^N (Y_p - \bar{Y})^2$$

In both formulas, all we're doing is summing over all of the observations in the sample. Most of the time we would just use the simpler Y_p notation: the equation using Y_p is clearly the simpler of the two. However, when doing an ANOVA it's important to keep track of which participants belong in which groups, and we need to use the Y_{ik} notation to do this.

14.2.2 From variances to sums of squares

Okay, now that we've got a good grasp on how the variance is calculated, let's define something called the **total sum of squares**, which is denoted SS_{tot} . This is very simple: instead of averaging the squared deviations, which is what we do when calculating the variance, we just add them up. So the formula for

the total sum of squares is almost identical to the formula for the variance:

$$SS_{tot} = \sum_{k=1}^G \sum_{i=1}^{N_k} (Y_{ik} - \bar{Y})^2$$

When we talk about analysing variances in the context of ANOVA, what we're really doing is working with the total sums of squares rather than the actual variance. One very nice thing about the total sum of squares is that we can break it up into two different kinds of variation. Firstly, we can talk about the **within-group sum of squares**, in which we look to see how different each individual person is from their own group mean:

$$SS_w = \sum_{k=1}^G \sum_{i=1}^{N_k} (Y_{ik} - \bar{Y}_k)^2$$

where \bar{Y}_k is a group mean. In our example, \bar{Y}_k would be the average mood change experienced by those people given the k -th drug. So, instead of comparing individuals to the average of all people in the experiment, we're only comparing them to those people in the same group. As a consequence, you'd expect the value of SS_w to be smaller than the total sum of squares, because it's completely ignoring any group differences – that is, the fact that the drugs (if they work) will have different effects on people's moods.

Next, we can define a third notion of variation which captures *only* the differences between groups. We do this by looking at the differences between the group means \bar{Y}_k and grand mean \bar{Y} . In order to quantify the extent of this variation, what we do is calculate the **between-group sum of squares**:

$$\begin{aligned} SS_b &= \sum_{k=1}^G \sum_{i=1}^{N_k} (\bar{Y}_k - \bar{Y})^2 \\ &= \sum_{k=1}^G N_k (\bar{Y}_k - \bar{Y})^2 \end{aligned}$$

It's not too difficult to show that the total variation among people in the experiment SS_{tot} is actually the sum of the differences between the groups SS_b and the variation inside the groups SS_w . That is:

$$SS_w + SS_b = SS_{tot}$$

Yay.

Okay, so what have we found out? We've discovered that the total variability associated with the outcome variable (SS_{tot}) can be mathematically carved up into the sum of “the variation due to the differences in the sample means for the different groups” (SS_b) plus “all the rest of the variation” (SS_w). How does that help me find out whether the groups have different population means? Um. Wait. Hold on a second... now that I think about it, this is *exactly* what we were looking for. If the null hypothesis is true, then you'd expect all the sample means to be pretty similar to each other, right? And that would imply that you'd expect SS_b to be really small, or at least you'd expect it to be a lot smaller than the “the variation associated with everything else”, SS_w . Hm. I detect a hypothesis test coming on...

14.2.3 From sums of squares to the F-test

As we saw in the last section, the *qualitative* idea behind ANOVA is to compare the two sums of squares values SS_b and SS_w to each other: if the between-group variation is SS_b is large relative to the within-group variation SS_w then we have reason to suspect that the population means for the different groups aren't identical to each other. In order to convert this into a workable hypothesis test, there's

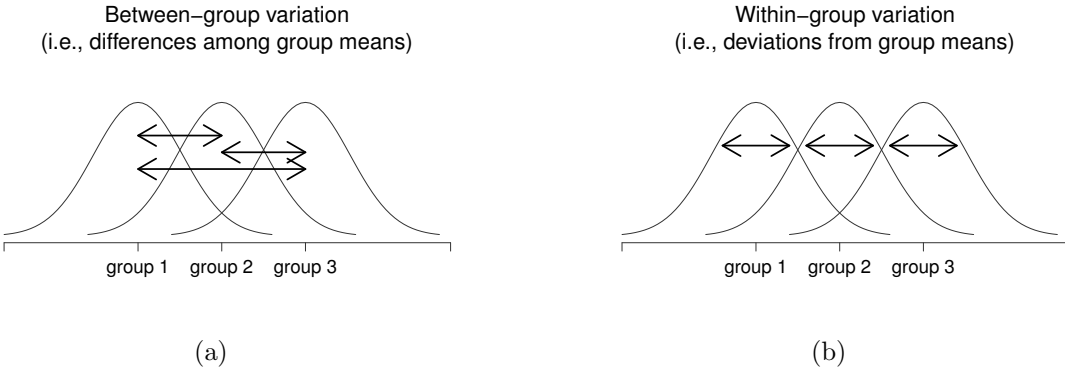


Figure 14.2: Graphical illustration of “between groups” variation (panel a) and “within groups” variation (panel b). On the left, the arrows show the differences in the group means; on the right, the arrows highlight the variability within each group.

a little bit of “fiddling around” needed. What I’ll do is first show you *what* we do to calculate our test statistic – which is called an ***F* ratio** – and then try to give you a feel for *why* we do it this way.

In order to convert our SS values into an *F*-ratio, the first thing we need to calculate is the **degrees of freedom** associated with the SS_b and SS_w values. As usual, the degrees of freedom corresponds to the number of unique “data points” that contribute to a particular calculation, minus the number of “constraints” that they need to satisfy. For the within-groups variability, what we’re calculating is the variation of the individual observations (N data points) around the group means (G constraints). In contrast, for the between groups variability, we’re interested in the variation of the group means (G data points) around the grand mean (1 constraint). Therefore, the degrees of freedom here are:

$$\begin{aligned} df_b &= G - 1 \\ df_w &= N - G \end{aligned}$$

Okay, that seems simple enough. What we do next is convert our summed squares value into a “mean squares” value, which we do by dividing by the degrees of freedom:

$$\begin{aligned} MS_b &= \frac{SS_b}{df_b} \\ MS_w &= \frac{SS_w}{df_w} \end{aligned}$$

Finally, we calculate the *F*-ratio by dividing the between-groups MS by the within-groups MS:

$$F = \frac{MS_b}{MS_w}$$

At a very general level, the intuition behind the *F* statistic is straightforward: bigger values of *F* means that the between-groups variation is large, relative to the within-groups variation. As a consequence, the larger the value of *F*, the more evidence we have against the null hypothesis. But how large does *F* have to be in order to actually *reject H_0* ? In order to understand this, you need a slightly deeper understanding of what ANOVA is and what the mean squares values actually are.

The next section discusses that in a bit of detail, but for readers that aren’t interested in the details of what the test is actually measuring, I’ll cut to the chase. In order to complete our hypothesis test, we

Table 14.1: All of the key quantities involved in an ANOVA, organised into a “standard” ANOVA table. The formulas for all quantities (except the p -value, which has a very ugly formula and would be nightmarishly hard to calculate without a computer) are shown.

	df	sum of squares	mean squares	F -statistic	p -value
between groups	$df_b = G - 1$	$SS_b = \sum_{k=1}^G N_k (\bar{Y}_k - \bar{Y})^2$	$MS_b = \frac{SS_b}{df_b}$	$F = \frac{MS_b}{MS_w}$	[complicated]
within groups	$df_w = N - G$	$SS_w = \sum_{k=1}^G \sum_{i=1}^{N_k} (Y_{ik} - \bar{Y}_k)^2$	$MS_w = \frac{SS_w}{df_w}$	-	-
.....					

need to know the sampling distribution for F if the null hypothesis is true. Not surprisingly, the sampling distribution for the F statistic under the null hypothesis is an F distribution. If you recall back to our discussion of the F distribution in Chapter 9, the F distribution has two parameters, corresponding to the two degrees of freedom involved: the first one df_1 is the between groups degrees of freedom df_b , and the second one df_2 is the within groups degrees of freedom df_w .

A summary of all the key quantities involved in a one-way ANOVA, including the formulas showing how they are calculated, is shown in Table 14.1.

14.2.4 The model for the data and the meaning of F

At a fundamental level, ANOVA is a competition between two different statistical models, H_0 and H_1 . When I described the null and alternative hypotheses at the start of the section, I was a little imprecise about what these models actually are. I’ll remedy that now, though you probably won’t like me for doing so. If you recall, our null hypothesis was that all of the group means are identical to one another. If so, then a natural way to think about the outcome variable Y_{ik} is to describe individual scores in terms of a single population mean μ , plus the deviation from that population mean. This deviation is usually denoted ϵ_{ik} and is traditionally called the *error* or *residual* associated with that observation. Be careful though: just like we saw with the word “significant”, the word “error” has a technical meaning in statistics that isn’t quite the same as its everyday English definition. In everyday language, “error” implies a mistake of some kind; in statistics, it doesn’t (or at least, not necessarily). With that in mind, the word “residual” is a better term than the word “error”. In statistics, both words mean “leftover variability”: that is, “stuff” that the model can’t explain. In any case, here’s what the null hypothesis looks like when we write it as a statistical model:

$$Y_{ik} = \mu + \epsilon_{ik}$$

where we make the *assumption* (discussed later) that the residual values ϵ_{ik} are normally distributed, with mean 0 and a standard deviation σ that is the same for all groups. To use the notation that we introduced in Chapter 9 we would write this assumption like this:

$$\epsilon_{ik} \sim \text{Normal}(0, \sigma^2)$$

What about the alternative hypothesis, H_1 ? The only difference between the null hypothesis and the alternative hypothesis is that we allow each group to have a different population mean. So, if we let μ_k denote the population mean for the k -th group in our experiment, then the statistical model corresponding to H_1 is:

$$Y_{ik} = \mu_k + \epsilon_{ik}$$

where, once again, we assume that the error terms are normally distributed with mean 0 and standard deviation σ . That is, the alternative hypothesis also assumes that $\epsilon \sim \text{Normal}(0, \sigma^2)$

Okay, now that we've described the statistical models underpinning H_0 and H_1 in more detail, it's now pretty straightforward to say what the mean square values are measuring, and what this means for the interpretation of F . I won't bore you with the proof of this, but it turns out that the within-groups mean square, MS_w , can be viewed as an estimator (in the technical sense: Chapter 10) of the error variance σ^2 . The between-groups mean square MS_b is also an estimator; but what it estimates is the error variance *plus* a quantity that depends on the true differences among the group means. If we call this quantity Q , then we can see that the F -statistic is basically²

$$F = \frac{\hat{Q} + \hat{\sigma}^2}{\hat{\sigma}^2}$$

where the true value $Q = 0$ if the null hypothesis is true, and $Q > 0$ if the alternative hypothesis is true (e.g. Hays, 1994, ch. 10). Therefore, at a bare minimum *the F value must be larger than 1* to have any chance of rejecting the null hypothesis. Note that this *doesn't* mean that it's impossible to get an F -value less than 1. What it means is that, if the null hypothesis is true the sampling distribution of the F ratio has a mean of 1,³ and so we need to see F -values larger than 1 in order to safely reject the null.

To be a bit more precise about the sampling distribution, notice that if the null hypothesis is true, both MS_b and MS_w are estimators of the variance of the residuals ϵ_{ik} . If those residuals are normally distributed, then you might suspect that the estimate of the variance of ϵ_{ik} is chi-square distributed... because (as discussed in Section 9.5) that's what a chi-square distribution *is*: it's what you get when you square a bunch of normally-distributed things and add them up. And since the F distribution is (again, by definition) what you get when you take the ratio between two things that are χ^2 distributed... we have our sampling distribution. Obviously, I'm glossing over a whole lot of stuff when I say this, but in broad terms, this really is where our sampling distribution comes from.

14.2.5 A worked example

{TODO} recheck all the numbers here

The previous discussion was fairly abstract, and a little on the technical side, so I think that at this point it might be useful to see a worked example. For that, let's go back to the clinical trial data that I introduced at the start of the chapter. The descriptive statistics that we calculated at the beginning tell us our group means: an average mood gain of 0.45 for the placebo, 0.72 for Anxifree, and 1.48 for Joyzepam. With that in mind, let's party like it's 1899⁴ and start doing some pencil and paper calculations. I'll only do this for the first 5 observations, because it's not bloody 1899 and I'm very lazy. Let's start by calculating SS_w , the within-group sums of squares. First, let's draw up a nice table to help

²In a later versions I'm intending to expand on this. But because I'm writing in a rush, and am already over my deadlines, I'll just briefly note that if you read ahead to Chapter 16 and look at how the "treatment effect" at level k of a factor is defined in terms of the α_k values (see Section 16.2), it turns out that Q refers to a weighted mean of the squared treatment effects, $Q = (\sum_{k=1}^G N_k \alpha_k^2)/(G - 1)$.

³Or, if we want to be sticklers for accuracy, $1 + \frac{2}{df_2 - 2}$.

⁴Or, to be precise, party like "it's 1899 and we've got no friends and nothing better to do with our time than do some calculations that wouldn't have made any sense in 1899 because ANOVA didn't exist until about the 1920s".

us with our calculations...

group <i>k</i>	outcome Y_{ik}
placebo	0.5
placebo	0.3
placebo	0.1
anxifree	0.6
anxifree	0.4

At this stage, the only thing I've included in the table is the raw data itself: that is, the grouping variable (i.e., `drug`) and outcome variable (i.e. `mood.gain`) for each person. Note that the outcome variable here corresponds to the Y_{ik} value in our equation previously. The next step in the calculation is to write down, for each person in the study, the corresponding group mean; that is, \bar{Y}_k . This is slightly repetitive, but not particularly difficult since we already calculated those group means when doing our descriptive statistics:

group <i>k</i>	outcome Y_{ik}	group mean \bar{Y}_k
placebo	0.5	0.45
placebo	0.3	0.45
placebo	0.1	0.45
anxifree	0.6	0.72
anxifree	0.4	0.72

Now that we've written those down, we need to calculate – again for every person – the deviation from the corresponding group mean. That is, we want to subtract $Y_{ik} - \bar{Y}_k$. After we've done that, we need to square everything. When we do that, here's what we get:

group <i>k</i>	outcome Y_{ik}	group mean \bar{Y}_k	dev. from group mean $Y_{ik} - \bar{Y}_k$	squared deviation $(Y_{ik} - \bar{Y}_k)^2$
placebo	0.5	0.45	0.05	0.0025
placebo	0.3	0.45	-0.15	0.0225
placebo	0.1	0.45	-0.35	0.1225
anxifree	0.6	0.72	-0.12	0.0136
anxifree	0.4	0.72	-0.32	0.1003

The last step is equally straightforward. In order to calculate the within-group sum of squares, we just add up the squared deviations across all observations:

$$\begin{aligned} SS_w &= 0.0025 + 0.0225 + 0.1225 + 0.0136 + 0.1003 \\ &= 0.2614 \end{aligned}$$

Of course, if we actually wanted to get the *right* answer, we'd need to do this for all 18 observations in the data set, not just the first five. We could continue with the pencil and paper calculations if we wanted to, but it's pretty tedious. Alternatively, it's not too hard to get R to do it. Here's how:

```
> outcome <- clin.trial$mood.gain
> group <- clin.trial$drug
> gp.means <- tapply(outcome, group, mean)
> gp.means <- gp.means[group]
> dev.from(gp.means) <- outcome - gp.means
> squared.devs <- dev.from(gp.means)^2
```

It might not be obvious from inspection what these commands are doing: as a general rule, the human brain seems to just shut down when faced with a big block of programming. However, I strongly suggest that – if you're like me and tend to find that the mere sight of this code makes you want to look away

and see if there's any beer left in the fridge or a game of footy on the telly – you take a moment and look closely at these commands one at a time. Every single one of these commands is something you've seen before somewhere else in the book. There's nothing novel about them (though I'll have to admit that the `tapply()` function takes a while to get a handle on), so if you're not quite sure how these commands work, this might be a good time to try playing around with them yourself, to try to get a sense of what's happening. On the other hand, if this does seem to make sense, then you won't be all that surprised at what happens when I wrap these variables in a data frame, and print it out...

```
> Y <- data.frame( group, outcome, gp.means,
+                   dev.from(gp.means, squared.devs )
> print(Y, digits = 2)
   group outcome gp.means dev.from(gp.means squared.devs
1 placebo      0.5     0.45       0.050      0.0025
2 placebo      0.3     0.45      -0.150      0.0225
3 placebo      0.1     0.45      -0.350      0.1225
4 anxifree     0.6     0.72      -0.117      0.0136
5 anxifree     0.4     0.72      -0.317      0.1003

BLAH BLAH BLAH

16 joyzepam    1.8     1.48       0.317      0.1003
17 joyzepam    1.3     1.48      -0.183      0.0336
18 joyzepam    1.4     1.48      -0.083      0.0069
```

If you compare this output to the contents of the table I've been constructing by hand, you can see that R has done exactly the same calculations that I was doing, and much faster too. So, if we want to finish the calculations of the within-group sum of squares in R, we just ask for the `sum()` of the `squared.devs` variable:

```
> SSw <- sum( squared.devs )
> print( SSw )
[1] 1.39
```

Obviously, this isn't the same as what I calculated, because R used all 18 observations. But if I'd typed `sum(squared.devs[1:5])` instead, it would have given the same answer that I got earlier.

Okay. Now that we've calculated the within groups variation, SS_w , it's time to turn our attention to the between-group sum of squares, SS_b . The calculations for this case are very similar. The main difference is that, instead of calculating the differences between an observation Y_{ik} and a group mean \bar{Y}_k for all of the observations, we calculate the differences between the group means \bar{Y}_k and the grand mean \bar{Y} (in this case 0.88) for all of the groups...

group <i>k</i>	group mean \bar{Y}_k	grand mean \bar{Y}	deviation $\bar{Y}_k - \bar{Y}$	squared deviations $(\bar{Y}_k - \bar{Y})^2$
placebo	0.45	0.88	-0.43	0.18
anxifree	0.72	0.88	-0.16	0.03
joyzepam	1.48	0.88	0.60	0.36

However, for the between group calculations we need to multiply each of these squared deviations by N_k , the number of observations in the group. We do this because every *observation* in the group (all N_k of them) is associated with a between group difference. So if there are six people in the placebo group, and the placebo group mean differs from the grand mean by 0.19, then the *total* between group variation

associated with these six people is $6 \times 0.16 = 1.14$. So we have to extend our little table of calculations...

group k	...	squared deviations $(\bar{Y}_k - \bar{Y})^2$	sample size N_k	weighted squared dev $N_k(\bar{Y}_k - \bar{Y})^2$
placebo	...	0.18	6	1.11
anxitfree	...	0.03	6	0.16
joyzepam	...	0.36	6	2.18

And so now our between group sum of squares is obtained by summing these “weighted squared deviations” over all three groups in the study:

$$\begin{aligned} SS_b &= 1.11 + 0.16 + 2.18 \\ &= 3.45 \end{aligned}$$

As you can see, the between group calculations are a lot shorter, so you probably wouldn’t usually want to bother using R as your calculator. However, if you *did* decide to do so, here’s one way you could do it:

```
> gp.means <- tapply(outcome, group, mean)
> grand.mean <- mean(outcome)
> dev.from.grand.mean <- gp.means - grand.mean
> squared.devs <- dev.from.grand.mean ^2
> gp.sizes <- tapply(outcome, group, length)
> wt.squared.devs <- gp.sizes * squared.devs
```

Again, I won’t actually try to explain this code line by line, but – just like last time – there’s nothing in there that we haven’t seen in several places elsewhere in the book, so I’ll leave it as an exercise for you to make sure you understand it. Once again, we can dump all our variables into a data frame so that we can print it out as a nice table:

```
> Y <- data.frame( gp.means, grand.mean, dev.from.grand.mean,
+                   squared.devs, gp.sizes, wt.squared.devs )
> print(Y, digits = 2)
      gp.means grand.mean dev.from.grand.mean squared.devs gp.sizes wt.squared.devs
placebo     0.45       0.88           -0.43      0.188      6        1.13
anxitfree   0.72       0.88           -0.17      0.028      6        0.17
joyzepam    1.48       0.88            0.60      0.360      6        2.16
```

Clearly, these are basically the same numbers that we got before. There are a few tiny differences, but that’s only because the hand-calculated versions have some small errors caused by the fact that I rounded all my numbers to 2 decimal places at each step in the calculations, whereas R only does it at the end (obviously, R’s version is more accurate). Anyway, here’s the R command showing the final step:

```
> SSb <- sum( wt.squared.devs )
> print( SSb )
[1] 3.453333
```

which is (ignoring the slight differences due to rounding error) the same answer that I got when doing things by hand.

Now that we’ve calculated our sums of squares values, SS_b and SS_w , the rest of the ANOVA is pretty painless. The next step is to calculate the degrees of freedom. Since we have $G = 3$ groups and $N = 18$ observations in total, our degrees of freedom can be calculated by simple subtraction:

$$\begin{aligned} df_b &= G - 1 = 2 \\ df_w &= N - G = 15 \end{aligned}$$

Next, since we've now calculated the values for the sums of squares and the degrees of freedom, for both the within-groups variability and the between-groups variability, we can obtain the mean square values by dividing one by the other:

$$\begin{aligned} \text{MS}_b &= \frac{\text{SS}_b}{\text{df}_b} = \frac{3.45}{2} = 1.73 \\ \text{MS}_w &= \frac{\text{SS}_w}{\text{df}_w} = \frac{1.39}{15} = 0.09 \end{aligned}$$

We're almost done. The mean square values can be used to calculate the F -value, which is the test statistic that we're interested in. We do this by dividing the between-groups MS value by the within-groups MS value.

$$F = \frac{\text{MS}_b}{\text{MS}_w} = \frac{1.73}{0.09} = 18.6$$

Woohooo! This is terribly exciting, yes? Now that we have our test statistic, the last step is to find out whether the test itself gives us a significant result. As discussed in Chapter 11, what we really *ought* to do is choose an α level (i.e., acceptable Type I error rate) ahead of time, construct our rejection region, etc etc. But in practice it's just easier to directly calculate the p -value. Back in the "old days", what we'd do is open up a statistics textbook or something and flick to the back section which would actually have a huge lookup table... that's how we'd "compute" our p -value, because it's too much effort to do it any other way. However, since we have access to R, I'll use the `pf()` function to do it instead. Now, remember that I explained earlier that the F -test is always one sided? And that we only reject the null hypothesis for very large F -values? That means we're only interested in the *upper tail* of the F -distribution. The command that you'd use here would be this...

```
> pf( 18.6, df1 = 2, df2 = 15, lower.tail = FALSE)
[1] 8.6727e-05
```

Therefore, our p -value comes to 0.0000867, or 8.67×10^{-5} in scientific notation. So, unless we're being *extremely* conservative about our Type I error rate, we're pretty much guaranteed to reject the null hypothesis.

At this point, we're basically done. Having completed our calculations, it's traditional to organise all these numbers into an ANOVA table like the one in Table 14.1. For our clinical trial data, the ANOVA table would look like this:

	df	sum of squares	mean squares	F -statistic	p -value
between groups	2	3.45	1.73	18.6	8.67×10^{-5}
within groups	15	1.39	0.09	-	-

These days, you'll probably never have much reason to want to construct one of these tables yourself, but you will find that almost all statistical software (R included) tends to organise the output of an ANOVA into a table like this, so it's a good idea to get used to reading them. However, although the software will output a full ANOVA table, there's almost never a good reason to include the whole table in your write up. A pretty standard way of reporting this result would be to write something like this:

One-way ANOVA showed a significant effect of drug on mood gain ($F(2, 15) = 18.6, p < .001$).

Sigh. So much work for one short sentence.

Running an ANOVA in R

I'm pretty sure I know what you're thinking after reading the last section, *especially* if you followed my advice and tried typing all the commands in yourself.... doing the ANOVA calculations yourself *sucks*. There's quite a lot of calculations that we needed to do along the way, and it would be tedious to have to do this over and over again every time you wanted to do an ANOVA. One possible solution to the problem would be to take all these calculations and turn them into some R functions yourself. You'd still have to do a lot of typing, but at least you'd only have to do it the one time: once you've created the functions, you can reuse them over and over again. However, writing your own functions is a lot of work, so this is kind of a last resort. Besides, it's much better if someone else does all the work for you...

14.3.1 Using the `aov()` function to specify your ANOVA

To make life easier for you, R provides a function called `aov()`, which – obviously – is an acronym of “Analysis Of Variance”.⁵ If you type `?aov` and have a look at the help documentation, you’ll see that there are several arguments to the `aov()` function, but the only two that we’re interested in are `formula` and `data`. As we’ve seen in a few places previously, the `formula` argument is what you use to specify the outcome variable and the grouping variable, and the `data` argument is what you use to specify the data frame that stores these variables. In other words, to do the same ANOVA that I laboriously calculated in the previous section, I’d use a command like this:

```
> aov( formula = mood.gain ~ drug, data = clin.trial )
```

Actually, that’s not *quite* the whole story, as you’ll see as soon as you look at the output from this command, which I’ve hidden for the moment in order to avoid confusing you. Before we go into specifics, I should point out that either of these commands will do the same thing:

```
> aov( clin.trial$mood.gain ~ clin.trial$drug )
> aov( mood.gain ~ drug, clin.trial )
```

In the first command, I didn’t specify a `data` set, and instead relied on the `$` operator to tell R how to find the variables. In the second command, I dropped the argument names, which is okay in this case because `formula` is the first argument to the `aov()` function, and `data` is the second one. Regardless of how I specify the ANOVA, I can assign the output of the `aov()` function to a variable, like this for example:

```
> my.anova <- aov( mood.gain ~ drug, clin.trial )
```

This is almost always a good thing to do, because there’s *lots* of useful things that we can do with the `my.anova` variable. So let’s assume that it’s this last command that I used to specify the ANOVA that I’m trying to run, and as a consequence I have this `my.anova` variable sitting in my workspace, waiting for me to do something with it...

14.3.2 Understanding what the `aov()` function produces

Now that we’ve seen how to use the `aov()` function to create `my.anova` we’d better have a look at what this variable actually is. The first thing to do is to check to see what class of variable we’ve created, since

⁵Actually, it *also* provides a function called `anova()`, but that works a bit differently, so let’s just ignore it for now.

it's kind of interesting in this case. When we do that...

```
> class( my.anova )
[1] "aov" "lm"
```

... we discover that `my.anova` actually has *two* classes! The first class tells us that it's an `aov` (analysis of variance) object, but the second tells us that it's *also* an `lm` (linear model) object. Later on, we'll see that this reflects a pretty deep statistical relationship between ANOVA and regression (Chapter 15) and it means that any function that exists in R for dealing with regressions can also be applied to `aov` objects, which is neat; but I'm getting ahead of myself. For now, I want to note that what we've created is an `aov` object, and to also make the point that `aov` objects are actually rather complicated beasts. I *won't* be trying to explain everything about them, since it's way beyond the scope of an introductory statistics subject, but to give you a tiny hint of some of the stuff that R stores inside an `aov` object, let's ask it to print out the `names()` of all the stored quantities...

```
> names( my.anova )
[1] "coefficients"   "residuals"      "effects"
[4] "rank"           "fitted.values"  "assign"
[7] "qr"             "df.residual"   "contrasts"
[10] "xlevels"        "call"          "terms"
[13] "model"
```

As we go through the rest of the book, I hope that a few of these will become a little more obvious to you, but right now that's going to look pretty damned opaque. That's okay. You don't need to know any of the details about it right now, and most of it you don't need at all... what you *do* need to understand is that the `aov()` function does a lot of calculations for you, not just the basic ones that I outlined in the previous sections. What this means is that it's generally a good idea to create a variable like `my.anova` that stores the output of the `aov()` function... because later on, you can use `my.anova` as an input to lots of other functions: those other functions can pull out bits and pieces from the `aov` object, and calculate various other things that you might need.

Right then. The simplest thing you can do with an `aov` object is to `print()` it out. When we do that, it shows us a few of the key quantities of interest:

```
> print( my.anova )
Call:
aov(formula = mood.gain ~ drug, data = clin.trial)

Terms:
            drug Residuals
Sum of Squares 3.4533    1.3917
Deg. of Freedom     2         15

Residual standard error: 0.30459
Estimated effects may be unbalanced
```

Specifically, it prints out a reminder of the command that you used when you called `aov()` in the first place, shows you the sums of squares values, the degrees of freedom, and a couple of other quantities that we're not really interested in right now. Notice, however, that R doesn't use the names "between-group" and "within-group". Instead, it tries to assign more meaningful names: in our particular example, the *between groups* variance corresponds to the effect that the `drug` has on the outcome variable; and the *within groups* variance corresponds to the "leftover" variability, so it calls that the `residuals`. If we compare these numbers to the numbers that I calculated by hand in Section 14.2.5, you can see that they're identical... the between groups sums of squares is $SS_b = 3.45$, the within groups sums of squares is $SS_w = 1.39$, and the degrees of freedom are 2 and 15 respectively.

14.3.3 Running the hypothesis tests for the ANOVA

Okay, so we've verified that `my.anova` seems to be storing a bunch of the numbers that we're looking for, but the `print()` function didn't quite give us the output that we really wanted. Where's the F -value? The p -value? These are the most important numbers in our hypothesis test, but the `print()` function doesn't provide them. To get those numbers, we need to use a different function. Instead of asking R to `print()` out the `aov` object, we should have asked for a `summary()` of it.⁶ When we do that...

```
> summary( my.anova )
   Df Sum Sq Mean Sq F value    Pr(>F)
drug      2    3.45   1.727    18.6 8.6e-05 ***
Residuals 15   1.39   0.093
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

... we get all of the key numbers that we calculated earlier. We get the sums of squares, the degrees of freedom, the mean squares, the F -statistic, and the p -value itself. These are all identical to the numbers that we calculated ourselves when doing it the long and tedious way, and it's even organised into the same kind of ANOVA table that I showed in Table 14.1, and then filled out by hand in Section 14.2.5. The only things that are even slightly different is that some of the row and column names are a bit different.

14.4

Effect size

There's a few different ways you could measure the effect size in an ANOVA, but the most commonly used measures are η^2 (**eta squared**) and partial η^2 . For a one way analysis of variance they're identical to each other, so for the moment I'll just explain η^2 . The definition of η^2 is actually really simple:

$$\eta^2 = \frac{SS_b}{SS_{tot}}$$

That's all it is. So when I look at the ANOVA table above, I see that $SS_b = 3.45$ and $SS_{tot} = 3.45 + 1.39 = 4.84$. Thus we get an η^2 value of

$$\eta^2 = \frac{3.45}{4.84} = 0.71$$

The interpretation of η^2 is equally straightforward: it refers to the proportion of the variability in the outcome variable (`mood.gain`) that can be explained in terms of the predictor (`drug`). A value of $\eta^2 = 0$ means that there is no relationship at all between the two, whereas a value of $\eta^2 = 1$ means that the relationship is perfect. Better yet, the η^2 value is very closely related to a squared correlation (i.e., r^2). So, if you're trying to figure out whether a particular value of η^2 is big or small, it's sometimes useful to remember that

$$\eta = \sqrt{\frac{SS_b}{SS_{tot}}}$$

can be interpreted as if it referred to the *magnitude* of a Pearson correlation. So in our drugs example, the η^2 value of .71 corresponds to an η value of $\sqrt{.71} = .84$. If we think about this as being equivalent to a correlation of about .84, we'd conclude that the relationship between `drug` and `mood.gain` is strong.

The core packages in R don't include any functions for calculating η^2 . However, it's pretty straightforward to calculate it directly from the numbers in the ANOVA table. In fact, since I've already got the `SSw` and `SSb` variables lying around from my earlier calculations, I can do this:

⁶It's worth noting that you can get the same result by using the command `anova(my.anova)`.

```

> SStot <- SSb + SSw      # total sums of squares
> eta.squared <- SSb / SStot # eta-squared value
> print( eta.squared )
[1] 0.71276

```

However, since it can be tedious to do this the long way (especially when we start running more complicated ANOVAs, such as those in Chapter 16) I've included an `etaSquared()` function in the `lsr` package which will do it for you. For now, the only argument you need to care about is `x`, which should be the `aov` object corresponding to your ANOVA. When we do this, what we get as output is this:

```

> etaSquared( x = my.anova )
      eta.sq eta.sq.part
drug 0.7127623  0.7127623

```

The output here shows two different numbers. The first one corresponds to the η^2 statistic, precisely as described above. The second one refers to “partial η^2 ”, which is a somewhat different measure of effect size that I’ll describe later. For the simple ANOVA that we’ve just run, they’re the same number. But this won’t always be true once we start running more complicated ANOVAs.

14.5

Multiple comparisons and post hoc tests

Any time you run an ANOVA with more than two groups, and you end up with a significant effect, the first thing you’ll probably want to ask is which groups are actually different from one another. In our drugs example, our null hypothesis was that all three drugs (placebo, Anxifree and Joyzepam) have the exact same effect on mood. But if you think about it, the null hypothesis is actually claiming *three* different things all at once here. Specifically, it claims that:

- Your competitor’s drug (Anxifree) is no better than a placebo (i.e., $\mu_A = \mu_P$)
- Your drug (Joyzepam) is no better than a placebo (i.e., $\mu_J = \mu_P$)
- Anxifree and Joyzepam are equally effective (i.e., $\mu_J = \mu_A$)

If any one of those three claims is false, then the null hypothesis is also false. So, now that we’ve rejected our null hypothesis, we’re thinking that *at least* one of those things isn’t true. But which ones? All three of these propositions are of interest: you certainly want to know if your new drug Joyzepam is better than a placebo, and it would be nice to know how well it stacks up against an existing commercial alternative (i.e., Anxifree). It would even be useful to check the performance of Anxifree against the placebo: even if Anxifree has already been extensively tested against placebos by other researchers, it can still be very useful to check that your study is producing similar results to earlier work.

When we characterise the null hypothesis in terms of these three distinct propositions, it becomes clear that there are eight possible “states of the world” that we need to distinguish between:

possibility:	is $\mu_P = \mu_A$?	is $\mu_P = \mu_J$?	is $\mu_A = \mu_J$?	which hypothesis?
1	✓	✓	✓	null
2	✓	✓		alternative
3	✓		✓	alternative
4	✓			alternative
5		✓	✓	alternative
6		✓		alternative
7			✓	alternative
8				alternative

By rejecting the null hypothesis, we've decided that we *don't* believe that #1 is the true state of the world. The next question to ask is, which of the other seven possibilities *do* we think is right? When faced with this situation, it's usually helps to look at the data. For instance, if we look at the plots in Figure 14.1, it's tempting to conclude that Joyzepam is better than the placebo and better than Anxifree, but there's no real difference between Anxifree and the placebo. However, if we want to get a clearer answer about this, it might help to run some tests.

14.5.1 Running “pairwise” t -tests

How might we go about solving our problem? Given that we've got three separate pairs of means (placebo versus Anxifree, placebo versus Joyzepam, and Anxifree versus Joyzepam) to compare, what we could do is run three separate t -tests and see what happens. There's a couple of ways that we could do this. One method would be to construct new variables corresponding the groups you want to compare (e.g., `anxitfree`, `placebo` and `joyzepam`), and then run a t -test on these new variables:

```
> t.test( anxitfree, placebo, var.equal = TRUE )    # Student t-test
```

or, you could use the `subset` argument in the `t.test()` function to select only those observations corresponding to one of the two groups we're interested in:

```
> t.test( formula = mood.gain ~ drug,
+         data = clin.trial,
+         subset = drug %in% c("placebo", "anxitfree"),
+         var.equal = TRUE
+     )
```

See Chapter 7 if you've forgotten how the `%in%` operator works. Regardless of which version we do, R will print out the results of the t -test, though I haven't included that output here. If we go on to do this for all possible pairs of variables, we can look to see which (if any) pairs of groups are significantly different to each other. This “lots of t -tests idea” isn't a bad strategy, though as we'll see later on there are some problems with it. However, for the moment our bigger problem is that it's a *pain* to have to type in such a long command over and over again: for instance, if your experiment has 10 groups, then you have to run 45 t -tests. That's way too much typing.

To help keep the typing to a minimum, R provides a function called `pairwise.t.test()` that automatically runs all of the t -tests for you. There are three arguments that you need to specify, the outcome variable `x`, the group variable `g`, and the `p.adjust.method` argument, which “adjusts” the p -value in one way or another. I'll explain p -value adjustment in a moment, but for now we can just set `p.adjust.method = "none"` since we're not doing any adjustments. For our example, here's what we do:

```
> pairwise.t.test( x = clin.trial$mood.gain,    # outcome variable
+                   g = clin.trial$drug,        # grouping variable
```

```

+           p.adjust.method = "none"      # which correction to use?
+ )

Pairwise comparisons using t tests with pooled SD

data: clin.trial$mood.gain and clin.trial$drug

placebo anxifree
anxifree 0.15021 -
joyzepam 3e-05   0.00056

P value adjustment method: none

```

One thing that bugs me slightly about the `pairwise.t.test()` function is that you can't just give it an `aov` object, and have it produce this output. After all, I went to all that trouble earlier of getting R to create the `my.anova` variable and – as we saw in Section 14.3.2 – R has actually stored enough information inside it that I should just be able to get it to run all the pairwise tests using `my.anova` as an input. To that end, I've included a `posthocPairwiseT()` function in the `lsr` package that lets you do this. The idea behind this function is that you can just input the `aov` object itself,⁷ and then get the pairwise tests as an output. As of the current writing, `posthocPairwiseT()` is actually just a simple way of calling `pairwise.t.test()` function, but you should be aware that I intend to make some changes to it later on. Here's an example:

```

> posthocPairwiseT( x = my.anova, p.adjust.method = "none" )

Pairwise comparisons using t tests with pooled SD

data: mood.gain and drug

placebo anxifree
anxifree 0.15021 -
joyzepam 3e-05   0.00056

P value adjustment method: none

```

In later versions, I plan to add more functionality (e.g., adjusted confidence intervals), but for now I think it's at least kind of useful. To see why, let's suppose you've run your ANOVA and stored the results in `my.anova`, and you're happy using the Holm correction (the default method in `pairwise.t.test()`, which I'll explain this in a moment). In that case, all you have to do is type this:

```
> posthocPairwiseT( my.anova )
```

and R will output the test results. Much more convenient, I think.

14.5.2 Corrections for multiple testing

In the previous section I hinted that there's a problem with just running lots and lots of t -tests. The concern is that when running these analyses, what we're doing is going on a “fishing expedition”: we're running lots and lots of tests without much theoretical guidance, in the hope that some of them come

⁷I should point out that there are other functions in R for running multiple comparisons, and at least one of them works this way: the `TukeyHSD()` function takes an `aov` object as its input, and outputs Tukey's “honestly significant difference” tests. I talk about Tukey's HSD in Chapter 16.

up significant. This kind of theory-free search for group differences is referred to as **post hoc analysis** (“post hoc” being Latin for “after this”).⁸

It’s okay to run post hoc analyses, but a lot of care is required. For instance, the analysis that I ran in the previous section is actually pretty dangerous: each *individual t-test* is designed to have a 5% Type I error rate (i.e., $\alpha = .05$), and I ran three of these tests. Imagine what would have happened if my ANOVA involved 10 different groups, and I had decided to run 45 “post hoc” *t*-tests to try to find out which ones were significantly different from each other, you’d expect 2 or 3 of them to come up significant *by chance alone*. As we saw in Chapter 11, the central organising principle behind null hypothesis testing is that we seek to control our Type I error rate, but now that I’m running lots of *t*-tests at once, in order to determine the source of my ANOVA results, my actual Type I error rate across this whole *family* of tests has gotten completely out of control.

The usual solution to this problem is to introduce an adjustment to the *p*-value, which aims to control the total error rate across the family of tests (see Shaffer, 1995). An adjustment of this form, which is usually (but not always) applied because one is doing post hoc analysis, is often referred to as a **correction for multiple comparisons**, though it is sometimes referred to as “simultaneous inference”. In any case, there are quite a few different ways of doing this adjustment. I’ll discuss a few of them in this section and in Section 16.8, but you should be aware that there are many other methods out there (see, e.g., Hsu, 1996).

14.5.3 Bonferroni corrections

The simplest of these adjustments is called the **Bonferroni correction** (Dunn, 1961), and it’s very very simple indeed. Suppose that my post hoc analysis consists of m separate tests, and I want to ensure that the total probability of making *any* Type I errors at all is at most α .⁹ If so, then the Bonferroni correction just says “multiply all your raw *p*-values by m ”. If we let p denote the original *p*-value, and let p'_j be the corrected value, then the Bonferroni correction tells that:

$$p' = m \times p$$

And therefore, if you’re using the Bonferroni correction, you would reject the null hypothesis if $p' < \alpha$. The logic behind this correction is very straightforward. We’re doing m different tests; so if we arrange it so that each test has a Type I error rate of at most α/m , then the *total* Type I error rate across these tests cannot be larger than α . That’s pretty simple, so much so that in the original paper, the author writes:

The method given here is so simple and so general that I am sure it must have been used before this. I do not find it, however, so can only conclude that perhaps its very simplicity has kept statisticians from realizing that it is a very good method in some situations (Dunn, 1961, pp 52-53)

To use the Bonferroni correction in R, you can use the `pairwise.t.test()` function,¹⁰ making sure that you set `p.adjust.method = "bonferroni"`. Alternatively, since the whole reason why we’re doing these pairwise tests in the first place is because we have an ANOVA that we’re trying to understand, it’s

⁸If you *do* have some theoretical basis for wanting to investigate some comparisons but not others, it’s a different story. In those circumstances you’re not really running “post hoc” analyses at all: you’re making “planned comparisons”. I do talk about this situation later in the book (Section 16.9), but for now I want to keep things simple.

⁹It’s worth noting in passing that not all adjustment methods try to do this. What I’ve described here is an approach for controlling “family wise Type I error rate”. However, there are other post hoc tests seek to control the “false discovery rate”, which is a somewhat different thing.

¹⁰There’s also a function called `p.adjust()` in which you can input a vector of raw *p*-values, and it will output a vector of adjusted *p*-values. This can be handy sometimes. I should also note that more advanced users may wish to consider using some of the tools provided by the `multcomp` package.

probably more convenient to use the `posthocPairwiseT()` function in the `lsr` package, since we can use `my.anova` as the input:

```
> posthocPairwiseT( my.anova, p.adjust.method = "bonferroni")

Pairwise comparisons using t tests with pooled SD

data: mood.gain and drug

placebo anxifree
anxifree 0.4506 -
joyzepam 9.1e-05 0.0017

P value adjustment method: bonferroni
```

If we compare these three p -values to those that we saw in the previous section when we made no adjustment at all, it is clear that the only thing that R has done is multiply them by 3.

14.5.4 Holm corrections

Although the Bonferroni correction is the simplest adjustment out there, it's not usually the best one to use. One method that is often used instead is the **Holm correction** (Holm, 1979). The idea behind the Holm correction is to pretend that you're doing the tests sequentially; starting with the smallest (raw) p -value and moving onto the largest one. For the j -th largest of the p -values, the adjustment is *either*

$$p'_j = j \times p_j$$

(i.e., the biggest p -value remains unchanged, the second biggest p -value is doubled, the third biggest p -value is tripled, and so on), or

$$p'_j = p'_{j+1}$$

whichever one is larger. This might sound a little confusing, so let's go through it a little more slowly. Here's what the Holm correction does. First, you sort all of your p -values in order, from smallest to largest. For the smallest p -value all you do is multiply it by m , and you're done. However, for all the other ones it's a two-stage process. For instance, when you move to the second smallest p value, you first multiply it by $m - 1$. If this produces a number that is bigger than the adjusted p -value that you got last time, then you keep it. But if it's smaller than the last one, then you copy the last p -value. To illustrate how this works, consider the table below, which shows the calculations of a Holm correction for a collection of five p -values:

raw p	rank j	$p \times j$	Holm p
.001	5	.005	.005
.005	4	.020	.020
.019	3	.057	.057
.022	2	.044	.057
.103	1	.103	.103

Hopefully that makes things clear.

Although it's a little harder to calculate, the Holm correction has some very nice properties: it's more powerful than Bonferroni (i.e., it has a lower Type II error rate), but – counterintuitive as it might seem – it has the *same* Type I error rate. As a consequence, in practice there's never any reason to use the simpler Bonferroni correction, since it is always outperformed by the slightly more elaborate

Holm correction. Because of this, the Holm correction is the default one used by `pairwise.t.test()` and `posthocPairwiseT()`. To run the Holm correction in R, you could specify `p.adjust.method = "Holm"` if you wanted to, but since it's the default you can just to do this:

```
> posthocPairwiseT( my.anova )

Pairwise comparisons using t tests with pooled SD

data: mood.gain and drug

placebo anxifree
anxifree 0.1502 -
joyzepam 9.1e-05 0.0011

P value adjustment method: holm
```

As you can see, the biggest p -value (corresponding to the comparison between Anxifree and the placebo) is unaltered: at a value of .15, it is exactly the same as the value we got originally when we applied no correction at all. In contrast, the smallest p -value (Joyzepam versus placebo) has been multiplied by three.

14.5.5 Writing up the post hoc test

Finally, having run the post hoc analysis to determine which groups are significantly different to one another, you might write up the result like this:

Post hoc tests (using the Holm correction to adjust p) indicated that Joyzepam produced a significantly larger mood change than both Anxifree ($p = .001$) and the placebo ($p = 9.1 \times 10^{-5}$). We found no evidence that Anxifree performed better than the placebo ($p = .15$).

Or, if you don't like the idea of reporting exact p -values, then you'd change those numbers to $p < .01$, $p < .001$ and $p > .05$ respectively. Either way, the key thing is that you indicate that you used Holm's correction to adjust the p -values. And of course, I'm assuming that elsewhere in the write up you've included the relevant descriptive statistics (i.e., the group means and standard deviations), since these p -values on their own aren't terribly informative.

14.6

Assumptions of one-way ANOVA

Like any statistical test, analysis of variance relies on some assumptions about the data. There are three key assumptions that you need to be aware of: *normality*, *homogeneity of variance* and *independence*. If you remember back to Section 14.2.4 – which I hope you at least skimmed even if you didn't read the whole thing – I described the statistical models underpinning ANOVA, which I wrote down like this:

$$\begin{aligned} H_0 : Y_{ik} &= \mu + \epsilon_{ik} \\ H_1 : Y_{ik} &= \mu_k + \epsilon_{ik} \end{aligned}$$

In these equations μ refers to a single, grand population mean which is the same for all groups, and μ_k is the population mean for the k -th group. Up to this point we've been mostly interested in whether our

data are best described in terms of a single grand mean (the null hypothesis) or in terms of different group-specific means (the alternative hypothesis). This makes sense, of course: that's actually the important research question! However, all of our testing procedures have – implicitly – relied on a specific assumption about the residuals, ϵ_{ik} , namely that

$$\epsilon_{ik} \sim \text{Normal}(0, \sigma^2)$$

None of the maths works properly without this bit. Or, to be precise, you can still do all the calculations, and you'll end up with an F -statistic, but you have no guarantee that this F -statistic actually measures what you think it's measuring, and so any conclusions that you might draw on the basis of the F test might be wrong.

So, how do we check whether this assumption about the residuals is accurate? Well, as I indicated above, there are three distinct claims buried in this one statement, and we'll consider them separately.

- **Normality.** The residuals are assumed to be normally distributed. As we saw in Section 13.7, we can assess this by looking at QQ plots or running a Shapiro-Wilk test. I'll talk about this in an ANOVA context in Section 14.9.
- **Homogeneity of variance.** Notice that we've only got the one value for the population standard deviation (i.e., σ), rather than allowing each group to have its own value (i.e., σ_k). This is referred to as the homogeneity of variance (sometimes called homoscedasticity) assumption. ANOVA assumes that the population standard deviation is the same for all groups. We'll talk about this extensively in Section 14.7.
- **Independence.** The independence assumption is a little trickier. What it basically means is that, knowing one residual tells you nothing about any other residual. All of the ϵ_{ik} values are assumed to have been generated without any “regard for” or “relationship to” any of the other ones. There's not an obvious or simple way to test for this, but there are some situations that are clear violations of this: for instance, if you have a repeated-measures design, where each participant in your study appears in more than one condition, then independence doesn't hold; there's a special relationship between some observations... namely those that correspond to the same person! When that happens, you need to use something like repeated measures ANOVA. I don't currently talk about repeated measures ANOVA in this book, but it will be included in later versions.

14.6.1 How robust is ANOVA?

One question that people often want to know the answer to is the extent to which you can trust the results of an ANOVA if the assumptions are violated. Or, to use the technical language, how **robust** is ANOVA to violations of the assumptions. Due to deadline constraints I don't have the time to discuss this topic. This is a topic I'll cover in some detail in a later version of the book.

14.7

Checking the homogeneity of variance assumption

There's more than one way to skin a cat, as the saying goes, and more than one way to test the homogeneity of variance assumption, too (though for some reason no-one made a saying out of that). The most commonly used test for this that I've seen in the literature is the **Levene test** (Levene, 1960), and the closely related **Brown-Forsythe test** (Brown & Forsythe, 1974), both of which I'll describe here. Alternatively, you could use the Bartlett test, which is implemented in R via the `bartlett.test()` function, but I'll leave it as an exercise for the reader to go check that one out if you're interested.

Levene's test is shockingly simple. Suppose we have our outcome variable Y_{ik} . All we do is define a new variable, which I'll call Z_{ik} , corresponding to the absolute deviation from the group mean:

$$Z_{ik} = |Y_{ik} - \bar{Y}_k|$$

Okay, what good does this do us? Well, let's take a moment to think about what Z_{ik} actually is, and what we're trying to test. The value of Z_{ik} is a measure of how the i -th observation in the k -th group deviates from its group mean. And our null hypothesis is that all groups have the same variance; that is, the same overall deviations from the group means! So, the null hypothesis in a Levene's test is that the population means of Z are identical for all groups. Hm. So what we need now is a statistical test of the null hypothesis that all group means are identical. Where have we seen that before? Oh right, that's what ANOVA is... and so all that the Levene's test does is run an ANOVA on the new variable Z_{ik} .

What about the Brown-Forsythe test? Does that do anything particularly different? Nope. The only change from the Levene's test is that it constructs the transformed variable Z in a slightly different way, using deviations from the group *medians* rather than deviations from the group *means*. That is, for the Brown-Forsythe test,

$$Z_{ik} = |Y_{ik} - \text{median}_k(Y)|$$

where $\text{median}_k(Y)$ is the median for group k . Regardless of whether you're doing the standard Levene test or the Brown-Forsythe test, the test statistic – which is sometimes denoted F , but sometimes written as W – is calculated in exactly the same way that the F -statistic for the regular ANOVA is calculated, just using a Z_{ik} rather than Y_{ik} . With that in mind, let's just move on and look at how to run the test in R.

14.7.1 Running the Levene's test in R

Okay, so how do we run the Levene test? Obviously, since the Levene test is just an ANOVA, it would be easy enough to manually create the transformed variable Z_{ik} and then use the `aov()` function to run an ANOVA on that. However, that's the tedious way to do it. A better way to do run your Levene's test is to use the `leveneTest()` function, which is in the `car` package. As usual, we first load the package

```
> library( car )
```

and now that we have, we can run our Levene test. The main argument that you need to specify is `y`, but you can do this in lots of different ways. Probably the simplest way to do it is actually input the original `aov` object. Since I've got the `my.anova` variable stored from my original ANOVA, I can just do this:

```
> leveneTest( my.anova )
Levene's Test for Homogeneity of Variance (center = median)
  Df F value Pr(>F)
group  2     1.47   0.26
      15
```

If we look at the output, we see that the test is non-significant ($F_{2,15} = 1.47, p = .26$), so it looks like the homogeneity of variance assumption is fine. Remember, although R reports the test statistic as an F -value, it could equally be called W , in which case you'd just write $W_{2,15} = 1.47$. Also, note the part of the output that says `center = median`. That's telling you that, by default, the `leveneTest()` function actually does the Brown-Forsythe test. If you want to use the mean instead, then you need to explicitly set the `center` argument, like this:

```
> leveneTest( y = my.anova, center = mean )
Levene's Test for Homogeneity of Variance (center = mean)
```

```
Df F value Pr(>F)
group  2    1.45   0.27
       15
```

That being said, in most cases it's probably best to stick to the default value, since the Brown-Forsythe test is a bit more robust than the original Levene test.

14.7.2 Additional comments

Two more quick comments before I move onto a different topic. Firstly, as mentioned above, there are other ways of calling the `leveneTest()` function. Although the vast majority of situations that call for a Levene test involve checking the assumptions of an ANOVA (in which case you probably have a variable like `my.anova` lying around), sometimes you might find yourself wanting to specify the variables directly. Two different ways that you can do this are shown below:

```
> leveneTest(y = mood.gain ~ drug, data = clin.trial) # y is a formula in this case
> leveneTest(y = clin.trial$mood.gain, group = clin.trial$drug) # y is the outcome
```

Secondly, I did mention that it's possible to run a Levene test just using the `aov()` function. I don't want to waste a lot of space on this, but just in case some readers are interested in seeing how this is done, here's the code that creates the new variables and runs an ANOVA. If you are interested, feel free to run this to verify that it produces the same answers as the Levene test (i.e., with `center = mean`):

```
> Y <- clin.trial $ mood.gain      # the original outcome variable, Y
> G <- clin.trial $ drug          # the grouping variable, G
> gp.mean <- tapply(Y, G, mean)   # calculate group means
> Ybar <- gp.mean[G]             # group mean associated with each obs
> Z <- abs(Y - Ybar)            # the transformed variable, Z
> summary( aov(Z ~ G) )         # run the ANOVA
```

That said, I don't imagine that many people will care about this. Nevertheless, it's nice to know that you could do it this way if you wanted to. And for those of you who do try it, I think it helps to demystify the test a little bit when you can see – with your own eyes – the way in which Levene's test relates to ANOVA.

14.8

Removing the homogeneity of variance assumption

In our example, the homogeneity of variance assumption turned out to be a pretty safe one: the Levene test came back non-significant, so we probably don't need to worry. However, in real life we aren't always that lucky. How do we save our ANOVA when the homogeneity of variance assumption is violated? If you recall from our discussion of *t*-tests, we've seen this problem before. The Student *t*-test assumes equal variances, so the solution was to use the Welch *t*-test, which does not. In fact, Welch (1951) also showed how we can solve this problem for ANOVA too (the **Welch one-way test**). It's implemented in R using the `oneway.test()` function. The arguments that we'll need for our example are:

- **formula**. This is the model formula, which (as usual) needs to specify the outcome variable on the left hand side and the grouping variable on the right hand side: i.e., something like `outcome ~ group`.
- **data**. Specifies the data frame containing the variables.

- `var.equal`. If this is `FALSE` (the default) a Welch one-way test is run. If it is `TRUE` then it just runs a regular ANOVA.

The function also has a `subset` argument that lets you analyse only some of the observations and a `na.action` argument that tells it how to handle missing data, but these aren't necessary for our purposes. So, to run the Welch one-way ANOVA for our example, we would do this:

```
> oneway.test(mood.gain ~ drug, data = clin.trial)

One-way analysis of means (not assuming equal variances)

data: mood.gain and drug
F = 26.322, num df = 2.000, denom df = 9.493, p-value = 0.000134
```

To understand what's happening here, let's compare these numbers to what we got earlier in Section 14.3 when we ran our original ANOVA. To save you the trouble of flicking back, here are those numbers again, this time calculated by setting `var.equal = TRUE` for the `oneway.test()` function:

```
> oneway.test(mood.gain ~ drug, data = clin.trial, var.equal = TRUE)

One-way analysis of means

data: mood.gain and drug
F = 18.611, num df = 2, denom df = 15, p-value = 8.646e-05
```

Okay, so originally our ANOVA gave us the result $F(2, 15) = 18.6$, whereas the Welch one-way test gave us $F(2, 9.49) = 26.32$. In other words, the Welch test has reduced the within-groups degrees of freedom from 15 to 9.49, and the F -value has increased from 18.6 to 26.32.

14.9

Checking the normality assumption

Testing the normality assumption is relatively straightforward. We covered most of what you need to know in Section 13.7. The only thing we really need to know how to do is pull out the residuals (i.e., the ϵ_{ik} values) so that we can draw our QQ plot and run our Shapiro-Wilk test. First, let's extract the residuals. R provides a function called `residuals()` that will do this for us. If we pass our `my.anova` to this function, it will return the residuals. So let's do that:

```
> my.anova.residuals <- residuals(object = my.anova) # extract the residuals
```

We can print them out too, though it's not exactly an edifying experience. In fact, given that I'm on the verge of putting *myself* to sleep just typing this, it might be a good idea to skip that step. Instead, let's draw some pictures and run ourselves a hypothesis test:

```
> hist(x = my.anova.residuals)           # plot a histogram (similar to Figure 14.3a)
> qqnorm(y = my.anova.residuals)         # draw a QQ plot (similar to Figure 14.3b)
> shapiro.test(x = my.anova.residuals)   # run Shapiro-Wilk test

Shapiro-Wilk normality test

data: my.anova.residuals
W = 0.9602, p-value = 0.6053
```

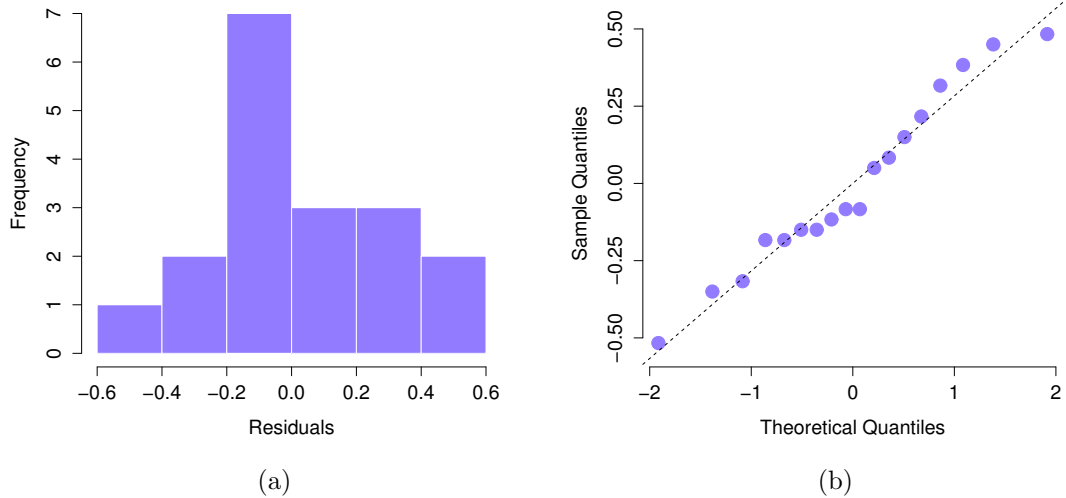


Figure 14.3: Histogram (panel a) and QQ plot (panel b) for the residuals of our ANOVA: both of these are in agreement with the Shapiro-Wilk test, and suggest that the residuals are normally distributed

The histogram and QQ plot are both shown in Figure 14.3, and both look pretty normal to me.¹¹ This is supported by the results of our Shapiro-Wilk test ($W = .96$, $p = .61$) which finds no indication that normality is violated.

14.10

Removing the normality assumption

Now that we've seen how to check for normality, we are led naturally to ask what we can do to address violations of normality. In the context of a one-way ANOVA, the easiest solution is probably to switch to a non-parametric test (i.e., one that doesn't rely on any particular assumption about the kind of distribution involved). We've seen non-parametric tests before, in Chapter 13: when you only have two groups, the Wilcoxon test provides the non-parametric alternative that you need. When you've got three or more groups, you can use the **Kruskal-Wallis rank sum test** (Kruskal & Wallis, 1952). So that's the test we'll talk about next.

14.10.1 The logic behind the Kruskal-Wallis test

The Kruskal-Wallis test is surprisingly similar to ANOVA, in some ways. In ANOVA, we started with Y_{ik} , the value of the outcome variable for the i th person in the k th group. For the Kruskal-Wallis test, what we'll do is rank order all of these Y_{ik} values, and conduct our analysis on the ranked data. So let's let R_{ik} refer to the ranking given to the i th member of the k th group. Now, let's calculate \bar{R}_k , the

¹¹Note that neither of these figures has been tidied up at all: if you want to create nicer looking graphs it's always a good idea to use the tools from Chapter 6 to help you draw cleaner looking images.

average rank given to observations in the k th group:

$$\bar{R}_k = \frac{1}{N_k} \sum_i R_{ik}$$

and let's also calculate \bar{R} , the grand mean rank:

$$\bar{R} = \frac{1}{N} \sum_i \sum_k R_{ik}$$

Now that we've done this, we can calculate the squared deviations from the grand mean rank \bar{R} . When we do this for the individual scores – i.e., if we calculate $(R_{ik} - \bar{R})^2$ – what we have is a “nonparametric” measure of how far the ik -th observation deviates from the grand mean rank. When we calculate the squared deviation of the group means from the grand means – i.e., if we calculate $(\bar{R}_k - \bar{R})^2$ – then what we have is a nonparametric measure of how much the *group* deviates from the grand mean rank. With this in mind, let's follow the same logic that we did with ANOVA, and define our *ranked* sums of squares measures in much the same way that we did earlier. First, we have our “total ranked sums of squares”:

$$RSS_{tot} = \sum_k \sum_i (R_{ik} - \bar{R})^2$$

and we can define the “between groups ranked sums of squares” like this:

$$\begin{aligned} RSS_b &= \sum_k \sum_i (\bar{R}_k - \bar{R})^2 \\ &= \sum_k N_k (\bar{R}_k - \bar{R})^2 \end{aligned}$$

So, if the null hypothesis is true and there are no true group differences at all, you'd expect the between group rank sums RSS_b to be very small, much smaller than the total rank sums RSS_{tot} . Qualitatively this is very much the same as what we found when we went about constructing the ANOVA F -statistic; but for technical reasons the Kruskal-Wallis test statistic, usually denoted K , is constructed in a slightly different way:

$$K = (N - 1) \times \frac{RSS_b}{RSS_{tot}}$$

and, if the null hypothesis is true, then the sampling distribution of K is *approximately* chi-square with $G - 1$ degrees of freedom (where G is the number of groups). The larger the value of K , the less consistent the data are with null hypothesis, so this is a one-sided test: we reject H_0 when K is sufficiently large.

14.10.2 Additional details

The description in the previous section illustrates the logic behind the Kruskal-Wallis test. At a conceptual level, this is the right way to think about how the test works. However, from a purely mathematical perspective it's needlessly complicated. I won't show you the derivation, but you can use a bit of algebraic jiggery-pokery¹² to show that the equation for K can be rewritten as

$$K = \frac{12}{N(N - 1)} \sum_k N_k \bar{R}_k^2 - 3(N + 1)$$

It's this last equation that you sometimes see given for K . This is way easier to calculate than the version I described in the previous section, it's just that it's totally meaningless to actual humans. It's probably best to think of K the way I described it earlier... as an analogue of ANOVA based on ranks. But keep

¹²A technical term.

in mind that the test statistic that gets calculated ends up with a rather different look to it than the one we used for our original ANOVA.

But wait, there's more! Dear lord, why is there always *more*? The story I've told so far is only actually true when there are no ties in the raw data. That is, if there are no two observations that have exactly the same value. If there *are* ties, then we have to introduce a correction factor to these calculations. At this point I'm assuming that even the most diligent reader has stopped caring (or at least formed the opinion that the tie-correction factor is something that doesn't require their immediate attention). So I'll very quickly tell you how it's calculated, and omit the tedious details about *why* it's done this way. Suppose we construct a frequency table for the raw data, and let f_j be the number of observations that have the j -th unique value. This might sound a bit abstract, so here's the R code showing a concrete example:

```
> f <- table( clin.trial$mood.gain )  # frequency table for mood gain
> print(f)   # we have some ties ...
0.1 0.2 0.3 0.4 0.5 0.6 0.8 0.9 1.1 1.2 1.3 1.4 1.7 1.8
1   1   2   1   1   2   1   1   1   1   2   2   1   1
```

Looking at this table, notice that the third entry in the frequency table has a value of 2. Since this corresponds to a `mood.gain` of 0.3, this table is telling us that two people's mood increased by 0.3. More to the point, note that we can say that `f[3]` has a value of `2`. Or, in the mathematical notation I introduced above, this is telling us that $f_3 = 2$. Yay. So, now that we know this, the tie correction factor (TCF) is:

$$\text{TCF} = 1 - \frac{\sum_j f_j^3 - f_j}{N^3 - N}$$

The tie-corrected value of the Kruskal-Wallis statistic obtained by dividing the value of K by this quantity: it is this tie-corrected version that R calculates. And at long last, we're actually finished with the theory of the Kruskal-Bloody-Wallis test. I'm sure you're all terribly relieved that I've cured you of the existential anxiety that naturally arises when you realise that you *don't* know how to calculate the tie-correction factor for the Kruskal-Wallis test. Right?

14.10.3 How to run the Kruskal-Wallis test in R

Despite the horror that we've gone through in trying to understand what the Kruskal-Wallis test actually does, it turns out that running the test is pretty painless, since R has a function called `kruskal.test()`. The function is pretty flexible, and allows you to input your data in a few different ways. Most of the time you'll have data like the `clin.trial` data set, in which you have your outcome variable `mood.gain`, and a grouping variable `drug`. If so, you can call the `kruskal.test()` function by specifying a formula, and a data frame:

```
> kruskal.test(mood.gain ~ drug, data = clin.trial)

Kruskal-Wallis rank sum test

data: mood.gain by drug
Kruskal-Wallis chi-squared = 12.076, df = 2, p-value = 0.002386
```

A second way of using the `kruskal.test()` function, which you probably won't have much reason to use, is to directly specify the outcome variable and the grouping variable as separate input arguments, `x` and `g`:

```
> kruskal.test(x = clin.trial$mood.gain, g = clin.trial$drug)
```

This isn't very interesting, since it's just plain easier to specify a formula. However, sometimes it can be useful to specify `x` as a list. What I mean is this. Suppose you actually had data as three separate variables, `placebo`, `anxitfree` and `joyzepam`. If that's the format that your data are in, then it's convenient to know that you can bundle all three together as a list:

```
> mood.gain <- list( placebo, joyzepam, anxitfree )
> kruskal.test( x = mood.gain )
```

And again, this would give you exactly the same results as the command we tried originally.

14.11

On the relationship between ANOVA and the Student *t* test

There's one last thing I want to point out before finishing. It's something that a lot of people find kind of surprising, but it's worth knowing about: an ANOVA with two groups is identical to the Student *t*-test. No, really. It's not just that they are similar, but they are actually equivalent in every meaningful way. I won't try to prove that this is always true, but I will show you a single concrete demonstration. Suppose that, instead of running an ANOVA on our `mood.gain ~ drug` model, let's instead do it using `therapy` as the predictor. If we run this ANOVA, here's what we get:

```
> summary( aov( mood.gain ~ therapy, data = clin.trial ) )
   Df Sum Sq Mean Sq F value Pr(>F)
therapy     1    0.47    0.467    1.71   0.21
Residuals  16   4.38   0.274
```

Overall, it looks like there's no significant effect here at all but, as we'll see in Chapter 16 this is actually a misleading answer! In any case, it's irrelevant to our current goals: our interest here is in the *F*-statistic, which is $F(1, 16) = 1.71$, and the *p*-value, which is .21. Since we only have two groups, I didn't actually need to resort to an ANOVA, I could have just decided to run a Student *t*-test. So let's see what happens when I do that:

```
> t.test( mood.gain ~ therapy, data = clin.trial, var.equal = TRUE )

Two Sample t-test

data: mood.gain by therapy
t = -1.3068, df = 16, p-value = 0.2098
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-0.84495  0.20051
sample estimates:
mean in group none  mean in group CBT
0.72222           1.04444
```

Curiously, the *p*-values are identical: once again we obtain a value of $p = .21$. But what about the test statistic? Having run a *t*-test instead of an ANOVA, we get a somewhat different answer, namely $t(16) = -1.3068$. However, there is a fairly straightforward relationship here. If we square the *t*-statistic

```
> 1.3068 ^ 2
[1] 1.7077
```

we get the F -statistic from before.

14.12

Summary

There's a fair bit covered in this chapter, but there's still a lot missing. Most obviously, I haven't yet discussed any analog of the paired samples t -test for more than two groups. There is a way of doing this, known as *repeated measures ANOVA*, which will appear in a later version of this book. I also haven't discussed how to run an ANOVA when you are interested in more than one grouping variable, but that will be discussed in a lot of detail in Chapter 16. In terms of what we have discussed, the key topics were:

- The basic logic behind how ANOVA works (Section 14.2) and how to run one in R(Section 14.3).
- How to compute an effect size for an ANOVA (Section 14.4)
- Post hoc analysis and corrections for multiple testing (Section 14.5).
- The assumptions made by ANOVA (Section 14.6).
- How to check the homogeneity of variance assumption (Section 14.7) and what to do if it is violated (Section 14.8).
- How to check the normality assumption (Section 14.9) and what to do if it is violated (Section 14.10).

15. Linear regression

The goal in this chapter is to introduce **linear regression**, the standard tool that statisticians rely on when analysing the relationship between interval scale predictors and interval scale outcomes. Stripped to its bare essentials, linear regression models are basically a slightly fancier version of the Pearson correlation (Section 5.7) though as we'll see, regression models are much more powerful tools.

15.1

What is a linear regression model?

Since the basic ideas in regression are closely tied to correlation, we'll return to the `parenthood.Rdata` file that we were using to illustrate how correlations work. Recall that, in this data set, we were trying to find out why Dan is so very grumpy all the time, and our working hypothesis was that I'm not getting enough sleep. We drew some scatterplots to help us examine the relationship between the amount of sleep I get, and my grumpiness the following day. The actual scatterplot that we draw is the one shown in Figure 15.1, and as we saw previously this corresponds to a correlation of $r = -.90$, but what we find ourselves secretly imagining is something that looks closer to Figure 15.2a. That is, we mentally draw a straight line through the middle of the data. In statistics, this line that we're drawing is called a **regression line**. Notice that – since we're not idiots – the regression line goes through the middle of the data. We don't find ourselves imagining anything like the rather silly plot shown in Figure 15.2b.

This is not highly surprising: the line that I've drawn in Figure 15.2b doesn't "fit" the data very well, so it doesn't make a lot of sense to propose it as a way of summarising the data, right? This is a very simple observation to make, but it turns out to be very powerful when we start trying to wrap just a little bit of maths around it. To do so, let's start with a refresher of some high school maths. The formula for a straight line is usually written like this:

$$y = mx + c$$

Or, at least, that's what it was when I went to high school all those years ago. The two *variables* are x and y , and we have two *coefficients*, m and c . The coefficient m represents the *slope* of the line, and the coefficient c represents the *y-intercept* of the line. Digging further back into our decaying memories of high school (sorry, for some of us high school was a long time ago), we remember that the intercept is interpreted as "the value of y that you get when $x = 0$ ". Similarly, a slope of m means that if you increase the x -value by 1 unit, then the y -value goes up by m units; a negative slope means that the y -value would go down rather than up. Ah yes, it's all coming back to me now.

Now that we've remembered that, it should come as no surprise to discover that we use the exact same formula to describe a regression line. If Y is the outcome variable (the DV) and X is the predictor

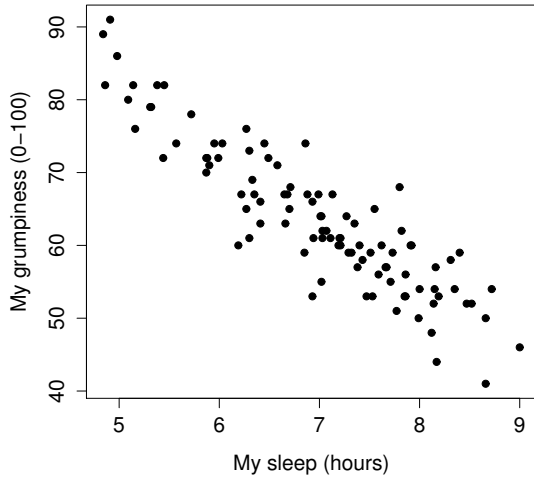


Figure 15.1: Scatterplot showing grumpiness as a function of hours slept.

variable (the IV), then the formula that describes our regression is written like this:

$$\hat{Y}_i = b_1 X_i + b_0$$

Hm. Looks like the same formula, but there's some extra frilly bits in this version. Let's make sure we understand them. Firstly, notice that I've written X_i and \hat{Y}_i rather than just plain old X and Y . This is because we want to remember that we're dealing with actual data. In this equation, X_i is the value of predictor variable for the i th observation (i.e., the number of hours of sleep that I got on day i of my little study), and \hat{Y}_i is the corresponding value of the outcome variable (i.e., my grumpiness on that day). And although I haven't said so explicitly in the equation, what we're assuming is that this formula works for all observations in the data set (i.e., for all i). Secondly, notice that I wrote \hat{Y}_i and not Y_i . This is because we want to make the distinction between the *actual data* Y_i , and the *estimate* \hat{Y}_i (i.e., the prediction that our regression line is making). Thirdly, I changed the letters used to describe the coefficients from m and c to b_1 and b_0 . That's just the way that statisticians like to refer to the coefficients in a regression model. I've no idea why they chose b , but that's what they did. In any case b_0 always refers to the intercept term, and b_1 refers to the slope.

Excellent, excellent. Next, I can't help but notice that – regardless of whether we're talking about the good regression line or the bad one – the data don't fall perfectly on the line. Or, to say it another way, the data Y_i are not identical to the predictions of the regression model \hat{Y}_i . Since statisticians love to attach letters, names and numbers to everything, let's refer to the difference between the model prediction and that actual data point as a *residual*, and we'll refer to it as ϵ_i .¹ Written using mathematics, the residuals are defined as:

$$\epsilon_i = Y_i - \hat{Y}_i$$

which in turn means that we can write down the complete linear regression model as:

$$Y_i = b_1 X_i + b_0 + \epsilon_i$$

¹The ϵ symbol is the Greek letter epsilon. It's traditional to use ϵ_i or e_i to denote a residual.

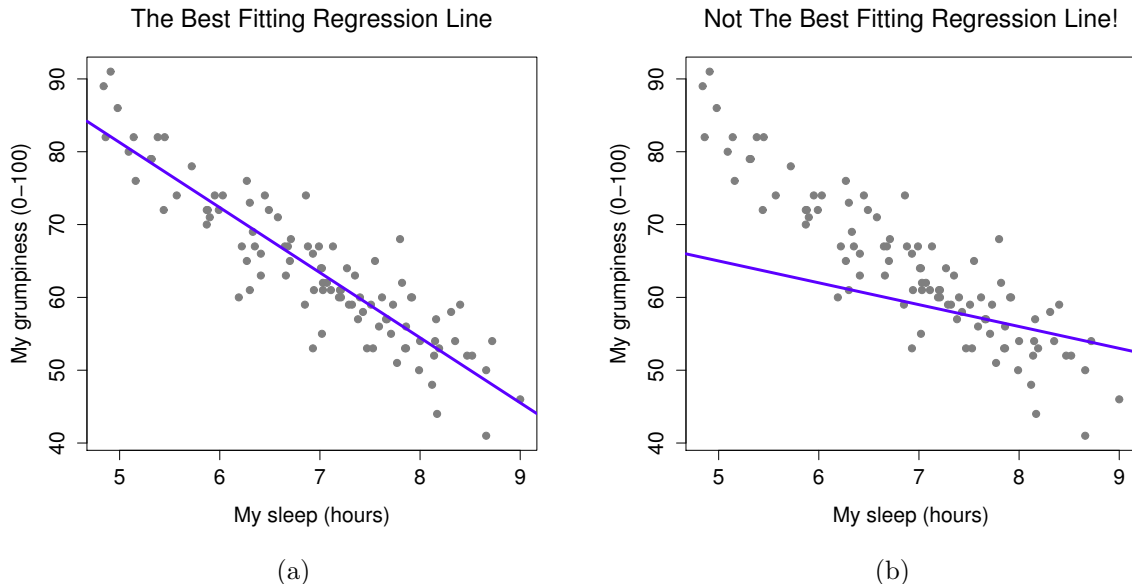


Figure 15.2: Panel a shows the sleep-grumpiness scatterplot from Figure 15.1 with the best fitting regression line drawn over the top. Not surprisingly, the line goes through the middle of the data. In contrast, panel b shows the same data, but with a very poor choice of regression line drawn over the top.

15.2

Estimating a linear regression model

Okay, now let's redraw our pictures, but this time I'll add some lines to show the size of the residual for all observations. When the regression line is good, our residuals (the lengths of the solid black lines) all look pretty small, as shown in Figure 15.3a, but when the regression line is a bad one, the residuals are a lot larger, as you can see from looking at Figure 15.3b. Hm. Maybe what we “want” in a regression model is *small* residuals. Yes, that does seem to make sense. In fact, I think I'll go so far as to say that the “best fitting” regression line is the one that has the smallest residuals. Or, better yet, since statisticians seem to like to take squares of everything why not say that ...

The estimated regression coefficients, \hat{b}_0 and \hat{b}_1 are those that minimise the sum of the squared residuals, which we could either write as $\sum_i(Y_i - \hat{Y}_i)^2$ or as $\sum_i \epsilon_i^2$.

Yes, yes that sounds even better. And since I've indented it like that, it probably means that this is the right answer. And since this is the right answer, it's probably worth making a note of the fact that our regression coefficients are *estimates* (we're trying to guess the parameters that describe a population!), which is why I've added the little hats, so that we get \hat{b}_0 and \hat{b}_1 rather than b_0 and b_1 . Finally, I should also note that – since there's actually more than one way to estimate a regression model – the more technical name for this estimation process is **ordinary least squares (OLS) regression**.

At this point, we now have a concrete definition for what counts as our “best” choice of regression coefficients, \hat{b}_0 and \hat{b}_1 . The natural question to ask next is, if our optimal regression coefficients are those that minimise the sum squared residuals, how do we *find* these wonderful numbers? The actual answer

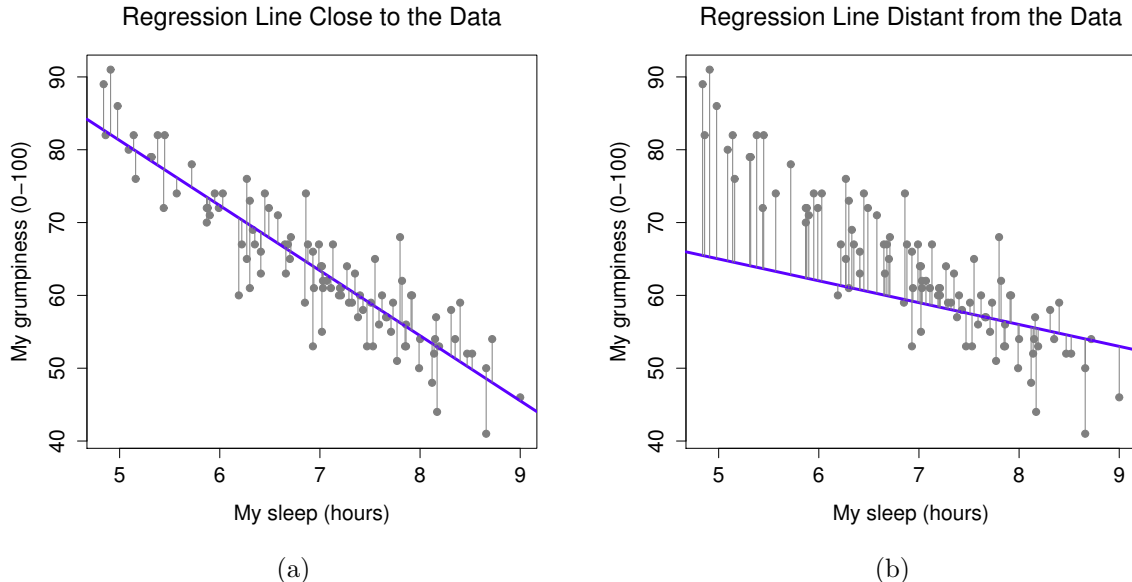


Figure 15.3: A depiction of the residuals associated with the best fitting regression line (panel a), and the residuals associated with a poor regression line (panel b). The residuals are much smaller for the good regression line. Again, this is no surprise given that the good line is the one that goes right through the middle of the data.

to this question is complicated, and it doesn't help you understand the logic of regression.² As a result, this time I'm going to let you off the hook. Instead of showing you how to do it the long and tedious way first, and then "revealing" the wonderful shortcut that R provides you with, let's cut straight to the chase... and use the `lm()` function (short for "linear model") to do all the heavy lifting.

15.2.1 Using the `lm()` function

The `lm()` function is a fairly complicated one: if you type `?lm`, the help files will reveal that there are a lot of arguments that you can specify, and most of them won't make a lot of sense to you. At this stage however, there's really only two of them that you care about, and as it turns out you've seen them before:

- **formula.** A formula that specifies the regression model. For the simple linear regression models that we've talked about so far, in which you have a single predictor variable as well as an intercept term, this formula is of the form `outcome ~ predictor`. However, more complicated formulas are allowed, and we'll discuss them later.

²Or at least, I'm assuming that it doesn't help most people. But on the off chance that someone reading this is a proper kung fu master of linear algebra (and to be fair, I always have a few of these people in my intro stats class), it *will* help *you* to know that the solution to the estimation problem turns out to be $\hat{\boldsymbol{b}} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$, where $\hat{\boldsymbol{b}}$ is a vector containing the estimated regression coefficients, \mathbf{X} is the "design matrix" that contains the predictor variables (plus an additional column containing all ones; strictly \mathbf{X} is a matrix of the regressors, but I haven't discussed the distinction yet), and \mathbf{y} is a vector containing the outcome variable. For everyone else, this isn't exactly helpful, and can be downright scary. However, since quite a few things in linear regression can be written in linear algebra terms, you'll see a bunch of footnotes like this one in this chapter. If you can follow the maths in them, great. If not, ignore it.

- **data**. The data frame containing the variables.

As we saw with `aov()` in Chapter 14, the output of the `lm()` function is a fairly complicated object, with quite a lot of technical information buried under the hood. Because this technical information is used by other functions, it's generally a good idea to create a variable that stores the results of your regression. With this in mind, to run my linear regression, the command I want to use is this:

```
> regression.1 <- lm( formula = dan.grump ~ dan.sleep,
+                      data = parenthood )
```

Note that I used `dan.grump ~ dan.sleep` as the formula: in the model that I'm trying to estimate, `dan.grump` is the *outcome* variable, and `dan.sleep` is the predictor variable. It's always a good idea to remember which one is which! Anyway, what this does is create an “`lm` object” (i.e., a variable whose class is “`lm`”) called `regression.1`. Let's have a look at what happens when we `print()` it out:

```
> print( regression.1 )

Call:
lm(formula = dan.grump ~ dan.sleep, data = parenthood)

Coefficients:
(Intercept)    dan.sleep
      125.956       -8.937
```

This looks promising. There's two separate pieces of information here. Firstly, R is politely reminding us what the command was that we used to specify the model in the first place, which can be helpful. More importantly from our perspective, however, is the second part, in which R gives us the intercept $\hat{b}_0 = 125.96$ and the slope $\hat{b}_1 = -8.94$. In other words, the best-fitting regression line that I plotted in Figure 15.2 has this formula:

$$\hat{Y}_i = -8.94 X_i + 125.96$$

15.2.2 Interpreting the estimated model

The most important thing to be able to understand is how to interpret these coefficients. Let's start with \hat{b}_1 , the slope. If we remember the definition of the slope, a regression coefficient of $\hat{b}_1 = -8.94$ means that if I increase X_i by 1, then I'm decreasing Y_i by 8.94. That is, each additional hour of sleep that I gain will improve my mood, reducing my grumpiness by 8.94 grumpiness points. What about the intercept? Well, since \hat{b}_0 corresponds to “the expected value of Y_i when X_i equals 0”, it's pretty straightforward. It implies that if I get zero hours of sleep ($X_i = 0$) then my grumpiness will go off the scale, to an insane value of ($Y_i = 125.96$). Best to be avoided, I think.

15.3

Multiple linear regression

The simple linear regression model that we've discussed up to this point assumes that there's a single predictor variable that you're interested in, in this case `dan.sleep`. In fact, up to this point, *every* statistical tool that we've talked about has assumed that your analysis uses one predictor variable and one outcome variable. However, in many (perhaps most) research projects you actually have multiple predictors that you want to examine. If so, it would be nice to be able to extend the linear regression

framework to be able to include multiple predictors. Perhaps some kind of **multiple regression** model would be in order?

Multiple regression is conceptually very simple. All we do is add more terms to our regression equation. Let's suppose that we've got two variables that we're interested in; perhaps we want to use both `dan.sleep` and `baby.sleep` to predict the `dan.grump` variable. As before, we let Y_i refer to my grumpiness on the i -th day. But now we have two X variables: the first corresponding to the amount of sleep I got and the second corresponding to the amount of sleep my son got. So we'll let X_{i1} refer to the hours I slept on the i -th day, and X_{i2} refers to the hours that the baby slept on that day. If so, then we can write our regression model like this:

$$Y_i = b_2 X_{i2} + b_1 X_{i1} + b_0 + \epsilon_i$$

As before, ϵ_i is the residual associated with the i -th observation, $\epsilon_i = Y_i - \hat{Y}_i$. In this model, we now have three coefficients that need to be estimated: b_0 is the intercept, b_1 is the coefficient associated with my sleep, and b_2 is the coefficient associated with my son's sleep. However, although the number of coefficients that need to be estimated has changed, the basic idea of how the estimation works is unchanged: our estimated coefficients \hat{b}_0 , \hat{b}_1 and \hat{b}_2 are those that minimise the sum squared residuals.

15.3.1 Doing it in R

Multiple regression in R is no different to simple regression: all we have to do is specify a more complicated `formula` when using the `lm()` function. For example, if we want to use both `dan.sleep` and `baby.sleep` as predictors in our attempt to explain why I'm so grumpy, then the formula we need is this:

```
dan.grump ~ dan.sleep + baby.sleep
```

Notice that, just like last time, I haven't explicitly included any reference to the intercept term in this formula; only the two predictor variables and the outcome. By default, the `lm()` function assumes that the model should include an intercept (though you can get rid of it if you want). In any case, I can create a new regression model – which I'll call `regression.2` – using the following command:

```
> regression.2 <- lm( formula = dan.grump ~ dan.sleep + baby.sleep,
+                         data = parenthood )
```

And just like last time, if we `print()` out this regression model we can see what the estimated regression coefficients are:

```
> print( regression.2 )

Call:
lm(formula = dan.grump ~ dan.sleep + baby.sleep, data = parenthood)

Coefficients:
(Intercept)    dan.sleep    baby.sleep
125.96557      -8.95025       0.01052
```

The coefficient associated with `dan.sleep` is quite large, suggesting that every hour of sleep I lose makes me a lot grumpier. However, the coefficient for `baby.sleep` is very small, suggesting that it doesn't really matter how much sleep my son gets; not really. What matters as far as my grumpiness goes is how much sleep *I* get. To get a sense of what this multiple regression model looks like, Figure 15.4 shows a 3D plot that plots all three variables, along with the regression model itself.

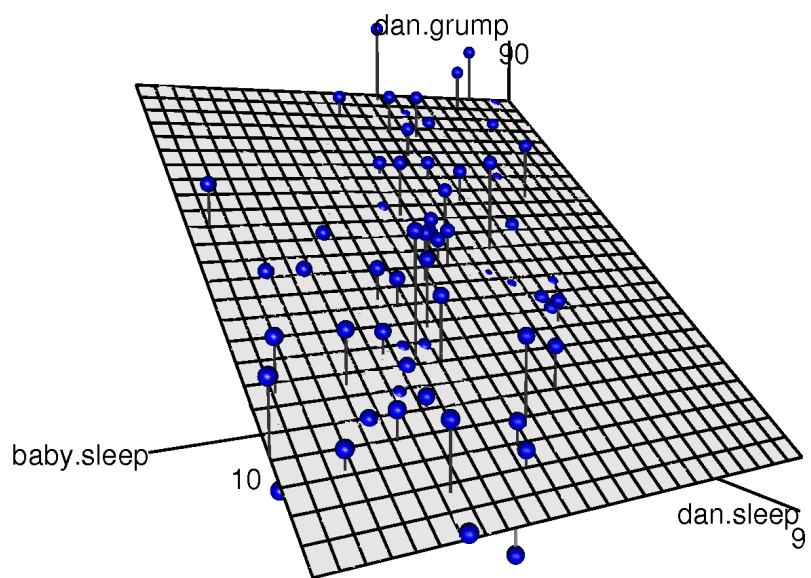


Figure 15.4: A 3D visualisation of a multiple regression model. There are two predictors in the model, `dan.sleep` and `baby.sleep`; the outcome variable is `dan.grump`. Together, these three variables form a 3D space: each observation (blue dots) is a point in this space. In much the same way that a simple linear regression model forms a line in 2D space, this multiple regression model forms a plane in 3D space. When we estimate the regression coefficients, what we're trying to do is find a plane that is as close to all the blue dots as possible. (This plot was drawn using the `scatter3d()` function in the `car` package, and it looked much nicer before it got butchered by the image conversion process that I used to get it into the book pdf)

15.3.2 Formula for the general case

The equation that I gave above shows you what a multiple regression model looks like when you include two predictors. Not surprisingly, then, if you want more than two predictors all you have to do is add more X terms and more b coefficients. In other words, if you have K predictor variables in the model then the regression equation looks like this:

$$Y_i = \left(\sum_{k=1}^K b_k X_{ik} \right) + b_0 + \epsilon_i$$

15.4

Quantifying the fit of the regression model

So we now know how to estimate the coefficients of a linear regression model. The problem is, we don't yet know if this regression model is any good. For example, the `regression.1` model *claims* that every hour of sleep will improve my mood by quite a lot, but it might just be rubbish. Remember, the regression model only produces a prediction \hat{Y}_i about what my mood is like: my actual mood is Y_i . If these two are very close, then the regression model has done a good job. If they are very different, then it has done a bad job.

15.4.1 The R^2 value

Once again, let's wrap a little bit of mathematics around this. Firstly, we've got the sum of the squared residuals:

$$SS_{res} = \sum_i (Y_i - \hat{Y}_i)^2$$

which we would hope to be pretty small. Specifically, what we'd like is for it to be very small in comparison to the total variability in the outcome variable,

$$SS_{tot} = \sum_i (Y_i - \bar{Y})^2$$

While we're here, let's calculate these values in R. Firstly, in order to make my R commands look a bit more similar to the mathematical equations, I'll create variables `X` and `Y`:

```
> X <- parenthood$dan.sleep # the predictor  
> Y <- parenthood$dan.grump # the outcome
```

Now that we've done this, let's calculate the \hat{Y} values and store them in a variable called `Y.pred`. For the simple model that uses only a single predictor, `regression.1`, we would do the following:

```
> Y.pred <- -8.94 * X + 125.97
```

Okay, now that we've got a variable which stores the regression model predictions for how grumpy I will be on any given day, let's calculate our sum of squared residuals. We would do that using the following command:

```
> SS.resid <- sum( (Y - Y.pred)^2 )  
> print( SS.resid )  
[1] 1838.722
```

Wonderful. A big number that doesn't mean very much. Still, let's forge boldly onwards anyway, and calculate the total sum of squares as well. That's also pretty simple:

```
> SS.tot <- sum( (Y - mean(Y))^2 )
> print( SS.tot )
[1] 9998.59
```

Hm. Well, it's a much bigger number than the last one, so this does suggest that our regression model was making good predictions. But it's not very interpretable.

Perhaps we can fix this. What we'd like to do is to convert these two fairly meaningless numbers into one number. A nice, interpretable number, which for no particular reason we'll call R^2 . What we would like is for the value of R^2 to be equal to 1 if the regression model makes no errors in predicting the data. In other words, if it turns out that the residual errors are zero – that is, if $SS_{res} = 0$ – then we expect $R^2 = 1$. Similarly, if the model is completely useless, we would like R^2 to be equal to 0. What do I mean by "useless"? Tempting as it is demand that the regression model move out of the house, cut its hair and get a real job, I'm probably going to have to pick a more practical definition: in this case, all I mean is that the residual sum of squares is no smaller than the total sum of squares, $SS_{res} = SS_{tot}$. Wait, why don't we do exactly that? The formula that provides us with our R^2 value is pretty simple to write down,

$$R^2 = 1 - \frac{SS_{res}}{SS_{tot}}$$

and equally simple to calculate in R:

```
> R.squared <- 1 - (SS.resid / SS.tot)
> print( R.squared )
[1] 0.8161018
```

The R^2 value, sometimes called the **coefficient of determination**³ has a simple interpretation: it is the *proportion* of the variance in the outcome variable that can be accounted for by the predictor. So in this case, the fact that we have obtained $R^2 = .816$ means that the predictor (`my.sleep`) explains 81.6% of the variance in the outcome (`my.grump`).

Naturally, you don't actually need to type in all these commands yourself if you want to obtain the R^2 value for your regression model. As we'll see later on in Section 15.5.3, all you need to do is use the `summary()` function. However, let's put that to one side for the moment. There's another property of R^2 that I want to point out.

15.4.2 The relationship between regression and correlation

At this point we can revisit my earlier claim that regression, in this very simple form that I've discussed so far, is basically the same thing as a correlation. Previously, we used the symbol r to denote a Pearson correlation. Might there be some relationship between the value of the correlation coefficient r and the R^2 value from linear regression? Of course there is: the squared correlation r^2 is identical to the R^2 value for a linear regression with only a single predictor. To illustrate this, here's the squared correlation:

```
> r <- cor(X, Y) # calculate the correlation
> print( r^2 )      # print the squared correlation
[1] 0.8161027
```

Yep, same number. In other words, running a Pearson correlation is more or less equivalent to running a linear regression model that uses only one predictor variable.

³And by "sometimes" I mean "almost never". In practice everyone just calls it "R-squared".

15.4.3 The adjusted R^2 value

One final thing to point out before moving on. It's quite common for people to report a slightly different measure of model performance, known as "adjusted R^2 ". The motivation behind calculating the adjusted R^2 value is the observation that adding more predictors into the model will *always* call the R^2 value to increase (or at least not decrease). The adjusted R^2 value introduces a slight change to the calculation, as follows. For a regression model with K predictors, fit to a data set containing N observations, the adjusted R^2 is:

$$\text{adj. } R^2 = 1 - \left(\frac{\text{SS}_{\text{res}}}{\text{SS}_{\text{tot}}} \times \frac{N - 1}{N - K - 1} \right)$$

This adjustment is an attempt to take the degrees of freedom into account. The big advantage of the adjusted R^2 value is that when you add more predictors to the model, the adjusted R^2 value will only increase if the new variables improve the model performance more than you'd expect by chance. The big disadvantage is that the adjusted R^2 value *can't* be interpreted in the elegant way that R^2 can. R^2 has a simple interpretation as the proportion of variance in the outcome variable that is explained by the regression model; to my knowledge, no equivalent interpretation exists for adjusted R^2 .

An obvious question then, is whether you should report R^2 or adjusted R^2 . This is probably a matter of personal preference. If you care more about interpretability, then R^2 is better. If you care more about correcting for bias, then adjusted R^2 is probably better. Speaking just for myself, I prefer R^2 : my feeling is that it's more important to be able to interpret your measure of model performance. Besides, as we'll see in Section 15.5, if you're worried that the improvement in R^2 that you get by adding a predictor is just due to chance and not because it's a better model, well, we've got hypothesis tests for that.

15.5

Hypothesis tests for regression models

So far we've talked about what a regression model is, how the coefficients of a regression model are estimated, and how we quantify the performance of the model (the last of these, incidentally, is basically our measure of effect size). The next thing we need to talk about is hypothesis tests. There are two different (but related) kinds of hypothesis tests that we need to talk about: those in which we test whether the regression model as a whole is performing significantly better than a null model; and those in which we test whether a particular regression coefficient is significantly different from zero.

At this point, you're probably groaning internally, thinking that I'm going to introduce a whole new collection of tests. You're probably sick of hypothesis tests by now, and don't want to learn any new ones. Me too. I'm so sick of hypothesis tests that I'm going to shamelessly reuse the F -test from Chapter 14 and the t -test from Chapter 13. In fact, all I'm going to do in this section is show you how those tests are imported wholesale into the regression framework.

15.5.1 Testing the model as a whole

Okay, suppose you've estimated your regression model. The first hypothesis test you might want to try is one in which the null hypothesis is that there is *no relationship* between the predictors and the outcome, and the alternative hypothesis is that *the data are distributed in exactly the way that the regression model predicts*. Formally, our "null model" corresponds to the fairly trivial "regression" model in which we include 0 predictors, and only include the intercept term b_0

$$H_0 : Y_i = b_0 + \epsilon_i$$

If our regression model has K predictors, the “alternative model” is described using the usual formula for a multiple regression model:

$$H_1 : Y_i = \left(\sum_{k=1}^K b_k X_{ik} \right) + b_0 + \epsilon_i$$

How can we test these two hypotheses against each other? The trick is to understand that just like we did with ANOVA, it’s possible to divide up the total variance SS_{tot} into the sum of the residual variance SS_{res} and the regression model variance SS_{mod} . I’ll skip over the technicalities, since we covered most of them in the ANOVA chapter, and just note that:

$$SS_{mod} = SS_{tot} - SS_{res}$$

And, just like we did with the ANOVA, we can convert the sums of squares into mean squares by dividing by the degrees of freedom.

$$MS_{mod} = \frac{SS_{mod}}{df_{mod}}$$

$$MS_{res} = \frac{SS_{res}}{df_{res}}$$

So, how many degrees of freedom do we have? As you might expect, the df associated with the model is closely tied to the number of predictors that we’ve included. In fact, it turns out that $df_{mod} = K$. For the residuals, the total degrees of freedom is $df_{res} = N - K - 1$.

Now that we’ve got our mean square values, you’re probably going to be entirely unsurprised (possibly even bored) to discover that we can calculate an F -statistic like this:

$$F = \frac{MS_{mod}}{MS_{res}}$$

and the degrees of freedom associated with this are K and $N - K - 1$. This F statistic has exactly the same interpretation as the one we introduced in Chapter 14. Large F values indicate that the null hypothesis is performing poorly in comparison to the alternative hypothesis. And since we already did some tedious “do it the long way” calculations back then, I won’t waste your time repeating them. In a moment I’ll show you how to do the test in R the easy way, but first, let’s have a look at the tests for the individual regression coefficients.

15.5.2 Tests for individual coefficients

The F -test that we’ve just introduced is useful for checking that the model as a whole is performing better than chance. This is important: if your regression model doesn’t produce a significant result for the F -test then you probably don’t have a very good regression model (or, quite possibly, you don’t have very good data). However, while failing this test is a pretty strong indicator that the model has problems, *passing* the test (i.e., rejecting the null) doesn’t imply that the model is good! Why is that, you might be wondering? The answer to that can be found by looking at the coefficients for the `regression.2` model:

```
> print( regression.2 )

Call:
lm(formula = dan.grump ~ dan.sleep + baby.sleep, data = parenthood)

Coefficients:
(Intercept)    dan.sleep    baby.sleep
 125.96557     -8.95025      0.01052
```

I can't help but notice that the estimated regression coefficient for the `baby.sleep` variable is tiny (0.01), relative to the value that we get for `dan.sleep` (-8.95). Given that these two variables are absolutely on the same scale (they're both measured in "hours slept"), I find this suspicious. In fact, I'm beginning to suspect that it's really only the amount of sleep that *I* get that matters in order to predict my grumpiness.

Once again, we can reuse a hypothesis test that we discussed earlier, this time the *t*-test. The test that we're interested has a null hypothesis that the true regression coefficient is zero ($b = 0$), which is to be tested against the alternative hypothesis that it isn't ($b \neq 0$). That is:

$$\begin{aligned} H_0 : b &= 0 \\ H_1 : b &\neq 0 \end{aligned}$$

How can we test this? Well, if the central limit theorem is kind to us, we might be able to guess that the sampling distribution of \hat{b} , the estimated regression coefficient, is a normal distribution with mean centred on b . What that would mean is that if the null hypothesis were true, then the sampling distribution of \hat{b} has mean zero and unknown standard deviation. Assuming that we can come up with a good estimate for the standard error of the regression coefficient, $\text{SE}(\hat{b})$, then we're in luck. That's *exactly* the situation for which we introduced the one-sample *t* way back in Chapter 13. So let's define a *t*-statistic like this,

$$t = \frac{\hat{b}}{\text{SE}(\hat{b})}$$

I'll skip over the reasons why, but our degrees of freedom in this case are $df = N - K - 1$. Irritatingly, the estimate of the standard error of the regression coefficient, $\text{SE}(\hat{b})$, is not as easy to calculate as the standard error of the mean that we used for the simpler *t*-tests in Chapter 13. In fact, the formula is somewhat ugly, and not terribly helpful to look at.⁴ For our purposes it's sufficient to point out that the standard error of the estimated regression coefficient depends on both the predictor and outcome variables, and is somewhat sensitive to violations of the homogeneity of variance assumption (discussed shortly).

In any case, this *t*-statistic can be interpreted in the same way as the *t*-statistics that we discussed in Chapter 13. Assuming that you have a two-sided alternative (i.e., you don't really care if $b > 0$ or $b < 0$), then it's the extreme values of *t* (i.e., a lot less than zero or a lot greater than zero) that suggest that you should reject the null hypothesis.

15.5.3 Running the hypothesis tests in R

To compute all of the quantities that we have talked about so far, all you need to do is ask for a `summary()` of your regression model. Since I've been using `regression.2` as my example, let's do that:

```
> summary( regression.2 )
```

The output that this command produces is pretty dense, but we've already discussed everything of interest in it, so what I'll do is go through it line by line. The first line reminds us of what the actual regression model is:

```
Call:
lm(formula = dan.grump ~ dan.sleep + baby.sleep, data = parenthood)
```

⁴For advanced readers only. The vector of residuals is $\epsilon = \mathbf{y} - \mathbf{X}\hat{b}$. For K predictors plus the intercept, the estimated residual variance is $\hat{\sigma}^2 = \epsilon'\epsilon/(N - K - 1)$. The estimated covariance matrix of the coefficients is $\hat{\sigma}^2(\mathbf{X}'\mathbf{X})^{-1}$, the main diagonal of which is $\text{SE}(\hat{b})$, our estimated standard errors.

You can see why this is handy, since it was a little while back when we actually created the `regression.2` model, and so it's nice to be reminded of what it was we were doing. The next part provides a quick summary of the residuals (i.e., the ϵ_i values),

```
Residuals:
    Min      1Q   Median      3Q     Max
-11.0345 -2.2198 -0.4016  2.6775 11.7496
```

which can be convenient as a quick and dirty check that the model is okay. Remember, we did assume that these residuals were normally distributed, with mean 0. In particular it's worth quickly checking to see if the median is close to zero, and to see if the first quartile is about the same size as the third quartile. If they look badly off, there's a good chance that the assumptions of regression are violated. These ones look pretty nice to me, so let's move on to the interesting stuff. The next part of the R output looks at the coefficients of the regression model:

```
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 125.96557   3.04095 41.423 <2e-16 ***
dan.sleep    -8.95025   0.55346 -16.172 <2e-16 ***
baby.sleep    0.01052   0.27106   0.039   0.969
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

Each row in this table refers to one of the coefficients in the regression model. The first row is the intercept term, and the later ones look at each of the predictors. The columns give you all of the relevant information. The first column is the actual estimate of b (e.g., 125.96 for the intercept, and -8.9 for the `dan.sleep` predictor). The second column is the standard error estimate $\hat{\sigma}_b$. The third column gives you the t -statistic, and it's worth noticing that in this table $t = \hat{b}/\text{SE}(\hat{b})$ every time. Finally, the fourth column gives you the actual p value for each of these tests.⁵ The only thing that the table itself doesn't list is the degrees of freedom used in the t -test, which is always $N - K - 1$ and is listed immediately below, in this line:

```
Residual standard error: 4.354 on 97 degrees of freedom
```

The value of $df = 97$ is equal to $N - K - 1$, so that's what we use for our t -tests. In the final part of the output we have the F -test and the R^2 values which assess the performance of the model as a whole

```
Residual standard error: 4.354 on 97 degrees of freedom
Multiple R-squared: 0.8161,          Adjusted R-squared: 0.8123
F-statistic: 215.2 on 2 and 97 DF,  p-value: < 2.2e-16
```

So in this case, the model performs significantly better than you'd expect by chance ($F(2, 97) = 215.2$, $p < .001$), which isn't all that surprising: the $R^2 = .812$ value indicate that the regression model accounts for 81.2% of the variability in the outcome measure. However, when we look back up at the t -tests for each of the individual coefficients, we have pretty strong evidence that the `baby.sleep` variable has no significant effect; all the work is being done by the `dan.sleep` variable. Taken together, these results suggest that `regression.2` is actually the wrong model for the data: you'd probably be better off dropping the `baby.sleep` predictor entirely. In other words, the `regression.1` model that we started with is the better model.

⁵Note that, although R has done multiple tests here, it hasn't done a Bonferroni correction or anything. These are standard one-sample t -tests with a two-sided alternative. If you want to make corrections for multiple tests, you need to do that yourself.

15.5.4 Hypothesis tests for correlations (a brief digression)

I don't want to spend too much time on this, but it's worth very briefly returning to the point I made earlier, that Pearson correlations are basically the same thing as linear regressions with only a single predictor added to the model. What this means is that the hypothesis tests that I just described in a regression context can also be applied to correlation coefficients. To see this, let's take a `summary()` of the `regression.1` model:

```
> summary( regression.1 )

Call:
lm(formula = dan.grump ~ dan.sleep, data = parenthood)

Residuals:
    Min      1Q  Median      3Q     Max 
-11.025 -2.213 -0.399  2.681 11.750 

Coefficients:
            Estimate Std. Error t value Pr(>|t|)    
(Intercept) 125.9563    3.0161   41.76 <2e-16 ***
dan.sleep    -8.9368    0.4285  -20.85 <2e-16 ***  
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1    1

Residual standard error: 4.332 on 98 degrees of freedom
Multiple R-squared:  0.8161,    Adjusted R-squared:  0.8142 
F-statistic: 434.9 on 1 and 98 DF,  p-value: < 2.2e-16
```

The important thing to note here is the t test associated with the predictor, in which we get a result of $t(98) = -20.85$, $p < .001$. Now let's compare this to the output of a different function, which goes by the name of `cor.test()`. As you might expect, this function runs a hypothesis test to see if the observed correlation between two variables is significantly different from 0. Let's have a look:

```
> cor.test( x = parenthood$dan.sleep, y = parenthood$dan.grump )

Pearson's product-moment correlation

data: parenthood$dan.sleep and parenthood$dan.grump
t = -20.8544, df = 98, p-value < 2.2e-16
alternative hypothesis: true correlation is not equal to 0
95 percent confidence interval:
-0.9340614 -0.8594714
sample estimates:
      cor
-0.903384
```

Again, the key thing to note is the line that reports the hypothesis test itself, which seems to be saying that $t(98) = -20.85$, $p < .001$. Hm. Looks like it's exactly the same test, doesn't it? And that's exactly what it is. The test for the significance of a correlation is identical to the t test that we run on a coefficient in a regression model.

Regarding regression coefficients

Before moving on to discuss the assumptions underlying linear regression and what you can do to check if they're being met, there's two more topics I want to briefly discuss, both of which relate to the regression coefficients. The first thing to talk about is calculating confidence intervals for the coefficients; after that, I'll discuss the somewhat murky question of how to determine which of predictor is most important.

15.6.1 Confidence intervals for the coefficients

Like any population parameter, the regression coefficients b cannot be estimated with complete precision from a sample of data; that's part of why we need hypothesis tests. Given this, it's quite useful to be able to report confidence intervals that capture our uncertainty about the true value of b . This is especially useful when the research question focuses heavily on an attempt to find out *how* strongly variable X is related to variable Y , since in those situations the interest is primarily in the regression weight b . Fortunately, confidence intervals for the regression weights can be constructed in the usual fashion,

$$\text{CI}(b) = \hat{b} \pm (t_{crit} \times \text{SE}(\hat{b}))$$

where $\text{SE}(\hat{b})$ is the standard error of the regression coefficient, and t_{crit} is the relevant critical value of the appropriate t distribution. For instance, if it's a 95% confidence interval that we want, then the critical value is the 97.5th quantile of a t distribution with $N - K - 1$ degrees of freedom. In other words, this is basically the same approach to calculating confidence intervals that we've used throughout. To do this in R we can use the `confint()` function. There arguments to this function are

- **object**. The regression model (`lm` object) for which confidence intervals are required.
- **parm**. A vector indicating which coefficients we should calculate intervals for. This can be either a vector of numbers or (more usefully) a character vector containing variable names. By default, all coefficients are included, so usually you don't bother specifying this argument.
- **level**. A number indicating the confidence level that should be used. As is usually the case, the default value is 0.95, so you wouldn't usually need to specify this argument.

So, suppose I want 99% confidence intervals for the coefficients in the `regression.2` model. I could do this using the following command:

```
> confint( object = regression.2,
+           level = .99
+ )
      0.5 %     99.5 %
(Intercept) 117.9755724 133.9555593
dan.sleep   -10.4044419 -7.4960575
baby.sleep   -0.7016868  0.7227357
```

Simple enough.

15.6.2 Calculating standardised regression coefficients

One more thing that you might want to do is to calculate “standardised” regression coefficients, often denoted β . The rationale behind standardised coefficients goes like this. In a lot of situations, your variables are on fundamentally different scales. Suppose, for example, my regression model aims to predict people’s IQ scores, using their educational attainment (number of years of education) and their income as predictors. Obviously, educational attainment and income are not on the same scales: the number of years of schooling can only vary by 10s of years, whereas income would vary by 10,000s of dollars (or more). The units of measurement have a big influence on the regression coefficients: the b coefficients only make sense when interpreted in light of the units, both of the predictor variables and the outcome variable. This makes it very difficult to compare the coefficients of different predictors. Yet there are situations where you really do want to make comparisons between different coefficients. Specifically, you might want some kind of standard measure of which predictors have the strongest relationship to the outcome. This is what **standardised coefficients** aim to do.

The basic idea is quite simple: the standardised coefficients are the coefficients that you would have obtained if you’d converted all the variables to z -scores before running the regression.⁶ The idea here is that, by converting all the predictors to z -scores, we they all go into the regression on the same scale, thereby removing the problem of having variables on different scales. Regardless of what the original variables were, a β value of 1 means that an increase in the predictor of 1 standard deviation will produce a corresponding 1 standard deviation increase in the outcome variable. Therefore, if variable A has a larger β value than variable B, it is deemed to have a stronger relationship with the outcome. Or at least that’s the idea: it’s worth being a little cautious here, since this does rely very heavily on the assumption that “a 1 standard deviation change” is fundamentally the same kind of thing for all variables. It’s not always obvious that this is true.

Leaving aside the interpretation issues, let’s look at how it’s calculated. What you could do is standardise all the variables yourself and then run a regression, but there’s a much simpler way to do it. As it turns out, the β coefficient for a predictor X and outcome Y has a very simple formula, namely

$$\beta_X = b_X \times \frac{\sigma_X}{\sigma_Y}$$

where σ_X is the standard deviation of the predictor, and σ_Y is the standard deviation of the outcome variable Y . This makes matters a lot simpler. To make things even simpler, the `lqr` package includes a function `standardCoefs()` that computes the β coefficients.

```
> standardCoefs( regression.2 )
      b          beta
dan.sleep -8.95024973 -0.90474809
baby.sleep  0.01052447  0.00217223
```

This clearly shows that the `dan.sleep` variable has a much stronger effect than the `baby.sleep` variable. However, this is a perfect example of a situation where it would probably make sense to use the original coefficients b rather than the standardised coefficients β . After all, my sleep and the baby’s sleep are *already* on the same scale: number of hours slept. Why complicate matters by converting these to z -scores?

⁶Strictly, you standardise all the *regressors*: that is, every “thing” that has a regression coefficient associated with it in the model. For the regression models that I’ve talked about so far, each predictor variable maps onto exactly one regressor, and vice versa. However, that’s not actually true in general: we’ll see some examples of this in Chapter 16. But for now, we don’t need to care too much about this distinction.

Assumptions of regression

The linear regression model that I've been discussing relies on several assumptions. In Section 15.8 we'll talk a lot more about how to check that these assumptions are being met, but first, let's have a look at each of them.

- *Normality.* Like half the models in statistics, standard linear regression relies on an assumption of normality. Specifically, it assumes that the *residuals* are normally distributed. It's actually okay if the predictors X and the outcome Y are non-normal, so long as the residuals ϵ are normal. See Section 15.8.3.
- *Linearity.* A pretty fundamental assumption of the linear regression model is that relationship between X and Y actually be linear! Regardless of whether it's a simple regression or a multiple regression, we assume that the relationships involved are linear. See Section 15.8.4.
- *Homogeneity of variance.* Strictly speaking, the regression model assumes that each residual ϵ_i is generated from a normal distribution with mean 0, and (more importantly for the current purposes) with a standard deviation σ that is the same for every single residual. In practice, it's impossible to test the assumption that every residual is identically distributed. Instead, what we care about is that the standard deviation of the residual is the same for all values of \hat{Y} , and (if we're being especially paranoid) all values of every predictor X in the model. See Section 15.8.5.
- *Uncorrelated predictors.* The idea here is that, in a multiple regression model, you don't want your predictors to be too strongly correlated with each other. This isn't "technically" an assumption of the regression model, but in practice it's required. Predictors that are too strongly correlated with each other (referred to as "collinearity") can cause problems when evaluating the model. See Section 15.8.6.
- *Residuals are independent of each other.* This is really just a "catch all" assumption, to the effect that "there's nothing else funny going on in the residuals". If there is something weird (e.g., the residuals all depend heavily on some other unmeasured variable) going on, it might screw things up.
- *No "bad" outliers.* Again, not actually a technical assumption of the model (or rather, it's sort of implied by all the others), but there is an implicit assumption that your regression model isn't being too strongly influenced by one or two anomalous data points; since this raises questions about the adequacy of the model, and the trustworthiness of the data in some cases. See Section 15.8.2.

Model checking

The main focus of this section is **regression diagnostics**, a term that refers to the art of checking that the assumptions of your regression model have been met, figuring out how to fix the model if the assumptions are violated, and generally to check that nothing "funny" is going on. I refer to this as the "art" of model checking with good reason: it's not easy, and while there are a lot of fairly standardised tools that you can use to diagnose and maybe even cure the problems that ail your model (if there are any, that is!), you really do need to exercise a certain amount of judgment when doing this. It's easy to get lost in all the details of checking this thing or that thing, and it's quite exhausting to try to remember

what all the different things are. This has the very nasty side effect that a lot of people get frustrated when trying to learn *all* the tools, so instead they decide not to do *any* model checking. This is a bit of a worry!

In this section, I describe several different things you can do to check that your regression model is doing what it's supposed to. It doesn't cover the full space of things you could do, but it's still much more detailed than what I see a lot of people doing in practice; and I don't usually cover all of this in my intro stats class myself. However, I do think it's important that you get a sense of what tools are at your disposal, so I'll try to introduce a bunch of them here. Finally, I should note that this section draws quite heavily from the [Fox and Weisberg \(2011\)](#) text, the book associated with the `car` package. The `car` package is notable for providing some excellent tools for regression diagnostics, and the book itself talks about them in an admirably clear fashion. I don't want to sound too gushy about it, but I do think that [Fox and Weisberg \(2011\)](#) is well worth reading.

15.8.1 Three kinds of residuals

The majority of regression diagnostics revolve around looking at the residuals, and by now you've probably formed a sufficiently pessimistic theory of statistics to be able to guess that – precisely *because* of the fact that we care a lot about the residuals – there are several different kinds of residual that we might consider. In particular, the following three kinds of residual are referred to in this section: “ordinary residuals”, “standardised residuals”, and “Studentised residuals”. There is a fourth kind that you'll see referred to in some of the Figures, and that's the “Pearson residual”: however, for the models that we're talking about in this chapter, the Pearson residual is identical to the ordinary residual.

The first and simplest kind of residuals that we care about are **ordinary residuals**. These are the actual, raw residuals that I've been talking about throughout this chapter. The ordinary residual is just the difference between the fitted value \hat{Y}_i and the observed value Y_i . I've been using the notation ϵ_i to refer to the i -th ordinary residual, and by gum I'm going to stick to it. With this in mind, we have the very simple equation

$$\epsilon_i = Y_i - \hat{Y}_i$$

This is of course what we saw earlier, and unless I specifically refer to some other kind of residual, this is the one I'm talking about. So there's nothing new here: I just wanted to repeat myself. In any case, you can get R to output a vector of ordinary residuals, you can use a command like this:

```
> residuals( object = regression.2 )
```

One drawback to using ordinary residuals is that they're always on a different scale, depending on what the outcome variable is and how good the regression model is. That is, Unless you've decided to run a regression model without an intercept term, the ordinary residuals will have mean 0; but the variance is different for every regression. In a lot of contexts, especially where you're only interested in the *pattern* of the residuals and not their actual values, it's convenient to estimate the **standardised residuals**, which are normalised in such a way as to have standard deviation 1. The way we calculate these is to divide the ordinary residual by an estimate of the (population) standard deviation of these residuals. For technical reasons, mumble mumble, the formula for this is:

$$\epsilon'_i = \frac{\epsilon_i}{\hat{\sigma}\sqrt{1 - h_i}}$$

where $\hat{\sigma}$ in this context is the estimated population standard deviation of the ordinary residuals, and h_i is the “hat value” of the i th observation. I haven't explained hat values to you yet (but have no fear,⁷ it's coming shortly), so this won't make a lot of sense. For now, it's enough to interpret the standardised

⁷Or have no hope, as the case may be.

residuals as if we'd converted the ordinary residuals to z -scores. In fact, that is more or less the truth, it's just that we're being a bit fancier. To get the standardised residuals, the command you want is this:

```
> rstandard( model = regression.2 )
```

Note that this function uses a different name for the input argument, but it's still just a linear regression object that the function wants to take as its input here.

The third kind of residuals are **Studentised residuals** (also called “jackknifed residuals”) and they're even fancier than standardised residuals. Again, the idea is to take the ordinary residual and divide it by some quantity in order to estimate some standardised notion of the residual, but the formula for doing the calculations this time is subtly different:

$$\epsilon_i^* = \frac{\epsilon_i}{\hat{\sigma}_{(-i)} \sqrt{1 - h_i}}$$

Notice that our estimate of the standard deviation here is written $\hat{\sigma}_{(-i)}$. What this corresponds to is the estimate of the residual standard deviation that you *would have obtained*, if you just deleted the i th observation from the data set. This sounds like the sort of thing that would be a nightmare to calculate, since it seems to be saying that you have to run N new regression models (even a modern computer might grumble a bit at that, especially if you've got a large data set). Fortunately, some terribly clever person has shown that this standard deviation estimate is actually given by the following equation:

$$\hat{\sigma}_{(-i)} = \hat{\sigma} \sqrt{\frac{N - K - 1 - \epsilon_i'^2}{N - K - 2}}$$

Isn't that a pip? Anyway, the command that you would use if you wanted to pull out the Studentised residuals for our regression model is

```
> rstudent( model = regression.2 )
```

Before moving on, I should point out that you don't often need to manually extract these residuals yourself, even though they are at the heart of almost all regression diagnostics. That is, the `residuals()`, `rstandard()` and `rstudent()` functions are all useful to *know* about, but most of the time the various functions that run the diagnostics will take care of these calculations for you. Even so, it's always nice to know how to actually get hold of these things yourself in case you ever need to do something non-standard.

15.8.2 Three kinds of anomalous data

One danger that you can run into with linear regression models is that your analysis might be disproportionately sensitive to a smallish number of “unusual” or “anomalous” observations. I discussed this idea previously in Section 6.5.2 in the context of discussing the outliers that get automatically identified by the `boxplot()` function, but this time we need to be much more precise. In the context of linear regression, there are three conceptually distinct ways in which an observation might be called “anomalous”. All three are interesting, but they have rather different implications for your analysis.

The first kind of unusual observation is an **outlier**. The definition of an outlier (in this context) is an observation that is very different from what the regression model predicts. An example is shown in Figure 15.5. In practice, we operationalise this concept by saying that an outlier is an observation that has a very large Studentised residual, ϵ_i^* . Outliers are interesting: a big outlier *might* correspond to junk data – e.g., the variables might have been entered incorrectly, or some other defect may be detectable. Note that you shouldn't throw an observation away just because it's an outlier. But the fact that it's an outlier is often a cue to look more closely at that case, and try to find out why it's so different.

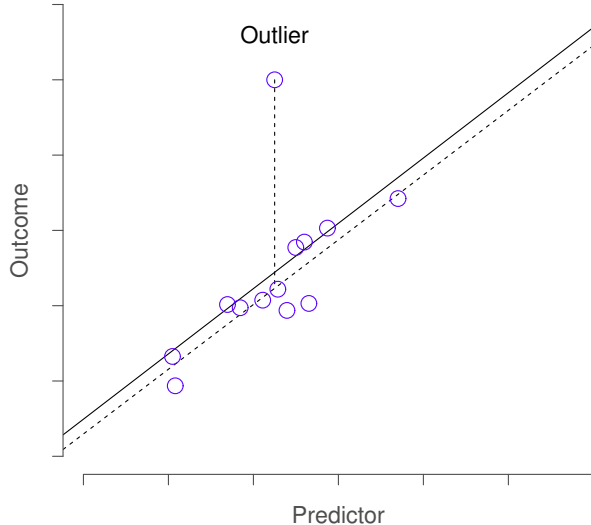


Figure 15.5: An illustration of outliers. The dotted lines plot the regression line that would have been estimated without the anomalous observation included, and the corresponding residual (i.e., the Studentised residual). The solid line shows the regression line with the anomalous observation included. The outlier has an unusual value on the outcome (y axis location) but not the predictor (x axis location), and lies a long way from the regression line.

The second way in which an observation can be unusual is if it has high **leverage**: this happens when the observation is very different from all the other observations. This doesn't necessarily have to correspond to a large residual: if the observation happens to be unusual on all variables in precisely the same way, it can actually lie very close to the regression line. An example of this is shown in Figure 15.6. The leverage of an observation is operationalised in terms of its *hat value*, usually written h_i . The formula for the hat value is rather complicated⁸ but its interpretation is not: h_i is a measure of the extent to which the i -th observation is “in control” of where the regression line ends up going. You can extract the hat values using the following command:

```
> hatvalues( model = regression.2 )
```

In general, if an observation lies far a way from the other ones in terms of the predictor variables, it will have a large hat value (as a rough guide, high leverage is when the hat value is more than 2-3 times the average; and note that the sum of the hat values is constrained to be equal to $K + 1$). High leverage points are also worth looking at in more detail, but they're much less likely to be a cause for concern unless they are also outliers.

⁸Again, for the linear algebra fanatics: the “hat matrix” is defined to be that matrix \mathbf{H} that converts the vector of observed values \mathbf{y} into a vector of fitted values $\hat{\mathbf{y}}$, such that $\hat{\mathbf{y}} = \mathbf{H}\mathbf{y}$. The name comes from the fact that this is the matrix that “puts a hat on \mathbf{y} ”. The hat *value* of the i -th observation is the i -th diagonal element of this matrix (so technically I should be writing it as h_{ii} rather than h_i). Oh, and in case you care, here's how it's calculated: $\mathbf{H} = \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'$. Pretty, isn't it?

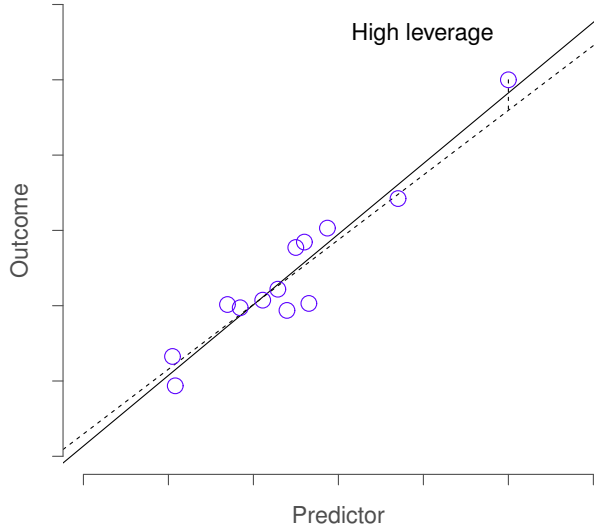


Figure 15.6: An illustration of high leverage points. The anomalous observation in this case is unusual both in terms of the predictor (x axis) and the outcome (y axis), but this unusualness is highly consistent with the pattern of correlations that exists among the other observations; as a consequence, the observation falls very close to the regression line and does not distort it.

This brings us to our third measure of unusualness, the **influence** of an observation. A high influence observation is an outlier that has high leverage. That is, it is an observation that is very different to all the other ones in some respect, and also lies a long way from the regression line. This is illustrated in Figure 15.7. Notice the contrast to the previous two figures: outliers don't move the regression line much, and neither do high leverage points. But something that is an outlier and has high leverage... that has a big effect on the regression line. That's why we call these points high influence; and it's why they're the biggest worry. We operationalise influence in terms of a measure known as **Cook's distance**,

$$D_i = \frac{\epsilon_i^{*2}}{K + 1} \times \frac{h_i}{1 - h_i}$$

Notice that this is a multiplication of something that measures the outlier-ness of the observation (the bit on the left), and something that measures the leverage of the observation (the bit on the right). In other words, in order to have a large Cook's distance, an observation must be a fairly substantial outlier *and* have high leverage. In a stunning turn of events, you can obtain these values using the following command:

```
> cooks.distance( model = regression.2 )
```

As a rough guide, Cook's distance greater than 1 is often considered large (that's what I typically use as a quick and dirty rule), though a quick scan of the internet and a few papers suggests that $4/N$ has also been suggested as a possible rule of thumb.

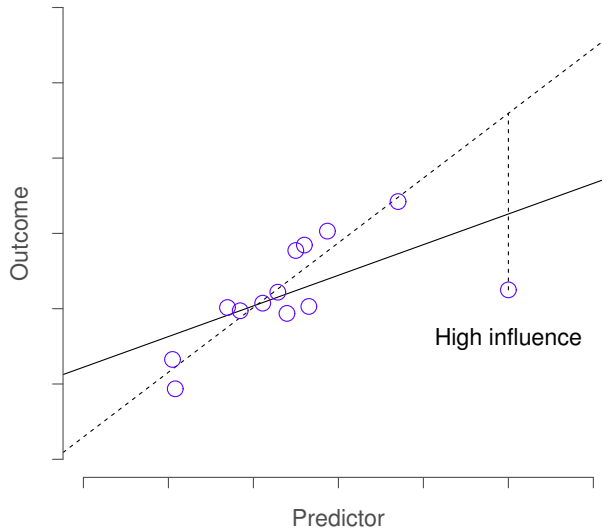


Figure 15.7: An illustration of high influence points. In this case, the anomalous observation is highly unusual on the predictor variable (x axis), and falls a long way from the regression line. As a consequence, the regression line is highly distorted, even though (in this case) the anomalous observation is entirely typical in terms of the outcome variable (y axis).

As hinted above, you don't usually need to make use of these functions, since you can have R automatically draw the critical plots.⁹ For the `regression.2` model, these are the plots showing Cook's distance (Figure 15.8) and the more detailed breakdown showing the scatter plot of the Studentised residual against leverage (Figure 15.9). To draw these, we can use the `plot()` function. When the main argument `x` to this function is a linear model object, it will draw one of six different plots, each of which is quite useful for doing regression diagnostics. You specify which one you want using the `which` argument (a number between 1 and 6). If you don't do this then R will draw all six. The two plots of interest to us in this context are generated using the following commands:

```
> plot(x = regression.2, which = 4) # Figure 15.8
> plot(x = regression.2, which = 5) # Figure 15.9
```

An obvious question to ask next is, if you do have large values of Cook's distance, what should you do? As always, there's no hard and fast rules. Probably the first thing to do is to try running the regression with that point excluded and see what happens to the model performance and to the regression coefficients. If they really are substantially different, it's time to start digging into your data set and your notes that you no doubt were scribbling as you ran your study; try to figure out *why* the point is so different. If you start to become convinced that this one data point is badly distorting your results, you might consider excluding it, but that's less than ideal unless you have a solid explanation for why this

⁹Though special mention should be made of the `influenceIndexPlot()` and `influencePlot()` functions in the `car` package. These produce somewhat more detailed pictures than the default plots that I've shown here. There's also an `outlierTest()` function that tests to see if any of the Studentised residuals are significantly larger than would be expected by chance.

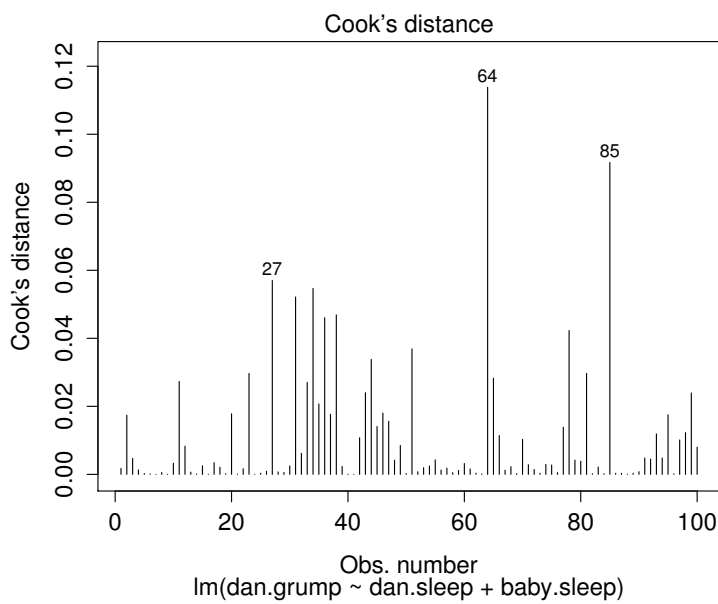


Figure 15.8: Cook's distance for every observation. This is one of the standard regression plots produced by the `plot()` function when the input is a linear regression object. It is obtained by setting `which=4`.

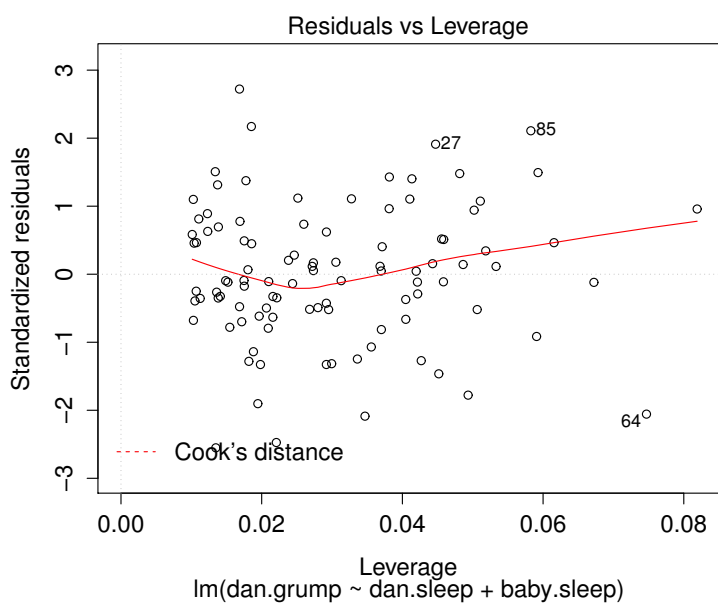


Figure 15.9: Residuals versus leverage. This is one of the standard regression plots produced by the `plot()` function when the input is a linear regression object. It is obtained by setting `which=5`.

particular case is qualitatively different from the others and therefore deserves to be handled separately.¹⁰ To give an example, let's delete the observation from day 64, the observation with the largest Cook's distance for the `regression.2` model. We can do this using the `subset` argument:

```
> lm( formula = dan.grump ~ dan.sleep + baby.sleep, # same formula
+     data = parenthood,          # same data frame...
+     subset = -64                # ...but observation 64 is deleted
+ )

Call:
lm(formula = dan.grump ~ dan.sleep + baby.sleep, data = parenthood,
subset = -64)

Coefficients:
(Intercept)    dan.sleep    baby.sleep
126.3553      -8.8283      -0.1319
```

As you can see, those regression coefficients have barely changed in comparison to the values we got earlier. In other words, we really don't have any problem as far as anomalous data are concerned.

15.8.3 Checking the normality of the residuals

Like many of the statistical tools we've discussed in this book, regression models rely on a normality assumption. In this case, we assume that the residuals are normally distributed. The tools for testing this aren't fundamentally different to those that we discussed earlier in Section 13.7. Firstly, I firmly believe that it never hurts to draw an old fashioned histogram. The command I use might be something like this:

```
> hist( x = residuals( regression.2 ),      # data are the residuals
+       xlab = "Value of residual",          # x-axis label
+       main = "",                          # no title
+       breaks = 20                         # lots of breaks
+ )
```

The resulting plot is shown in Figure 15.10, and as you can see the plot looks pretty damn close to normal, almost unnaturally so. I could also run a Shapiro-Wilk test to check, using the `shapiro.test()` function; the W value of .99, at this sample size, is non-significant ($p = .84$), again suggesting that the normality assumption isn't in any danger here. As a third measure, we might also want to draw a QQ-plot using the `qqnorm()` function. The QQ plot is an excellent one to draw, and so you might not be surprised to discover that it's one of the regression plots that we can produce using the `plot()` function:

```
> plot( x = regression.2, which = 2 )    # Figure 15.11
```

The output is shown in Figure 15.11, showing the standardised residuals plotted as a function of their theoretical quantiles according to the regression model. The fact that the output appends the model specification to the picture is nice.

15.8.4 Checking the linearity of the relationship

The third thing we might want to test is the linearity of the relationships between the predictors and the outcomes. There's a few different things that you might want to do in order to check this. Firstly, it

¹⁰An alternative is to run a “robust regression”; I'll discuss robust regression in a later version of this book.

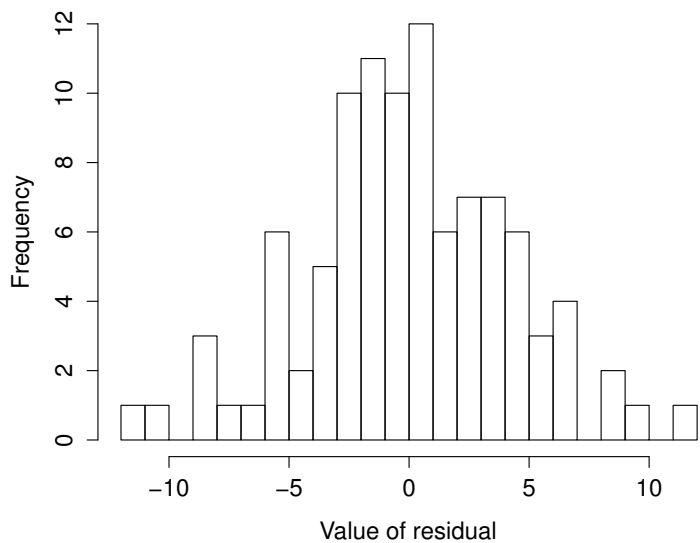


Figure 15.10: A histogram of the (ordinary) residuals in the `regression.2` model. These residuals look very close to being normally distributed, much moreso than is typically seen with real data. This shouldn't surprise you... they aren't real data, and they aren't real residuals!

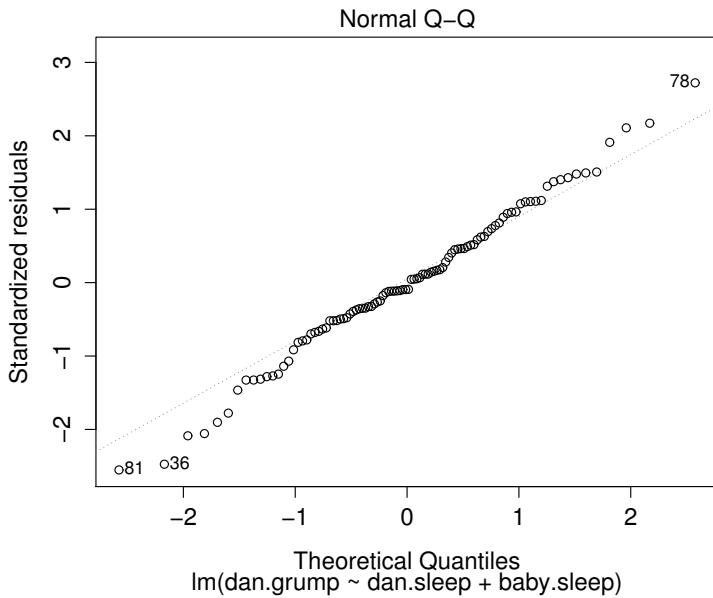


Figure 15.11: Plot of the theoretical quantiles according to the model, against the quantiles of the standardised residuals. This is one of the standard regression plots produced by the `plot()` function when the input is a linear regression object. It is obtained by setting `which=2`.

never hurts to just plot the relationship between the fitted values \hat{Y}_i and the observed values Y_i for the outcome variable, as illustrated in Figure 15.12. To draw this we could use the `fitted.values()` function to extract the \hat{Y}_i values in much the same way that we used the `residuals()` function to extract the ϵ_i values. So the commands to draw this figure might look like this:

```
> yhat.2 <- fitted.values( object = regression.2 )
> plot( x = yhat.2,
+       y = parenthood$dan.grump,
+       xlab = "Fitted Values",
+       ylab = "Observed Values"
+ )
```

One of the reasons I like to draw these plots is that they give you a kind of “big picture view”. If this plot looks approximately linear, then we’re probably not doing too badly (though that’s not to say that there aren’t problems). However, if you can see big departures from linearity here, then it strongly suggests that you need to make some changes.

In any case, in order to get a more detailed picture it’s often more informative to look at the relationship between the fitted values and the residuals themselves. Again, we could draw this plot using low level commands, but there’s an easier way. Just `plot()` the regression model, and select `which = 1`:

```
> plot(x = regression.2, which = 1) # Figure 15.13
```

The output is shown in Figure 15.13. As you can see, not only does it draw the scatterplot showing the fitted value against the residuals, it also plots a line through the data that shows the relationship between the two. Ideally, this should be a straight, perfectly horizontal line. There’s some hint of curvature here,

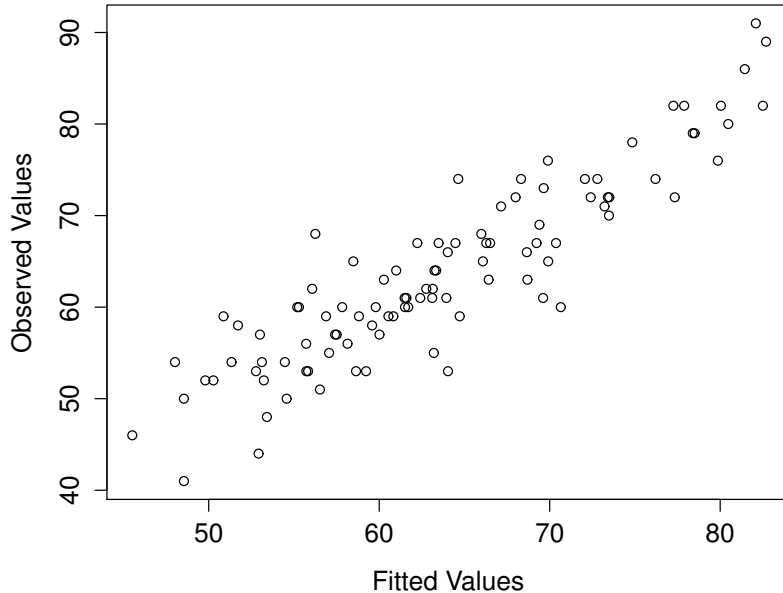


Figure 15.12: Plot of the fitted values against the observed values of the outcome variable. A straight line is what we're hoping to see here. This looks pretty good, suggesting that there's nothing grossly wrong, but there could be hidden subtle issues.

but it's not clear whether or not we be concerned.

A somewhat more advanced version of the same plot is produced by the `residualPlots()` function in the `car` package. This function not only draws plots comparing the fitted values to the residuals, it does so for each individual predictor. The command is

```
> residualPlots( model = regression.2 ) # Figure 15.14
```

and the resulting plots are shown in Figure 15.14. Note that this function also reports the results of a bunch of **curvature tests**. For a predictor variable X in some regression model, this test is equivalent to adding a new predictor to the model corresponding to X^2 , and running the t -test on the b coefficient associated with this new predictor. If it comes up significant, it implies that there is some nonlinear relationship between the variable and the residuals. For what it's worth, here's what you get for the `regression.2` model:

	Test stat	Pr(> t)
dan.sleep	2.160	0.033
baby.sleep	-0.545	0.587
Tukey test	2.162	0.031

The third line here is the **Tukey test**, which is basically the same test, except that instead of squaring one of the predictors and adding it to the model, you square the fitted-value. In any case, the fact that the curvature tests have come up significant is hinting that the curvature that we can see in Figures 15.13

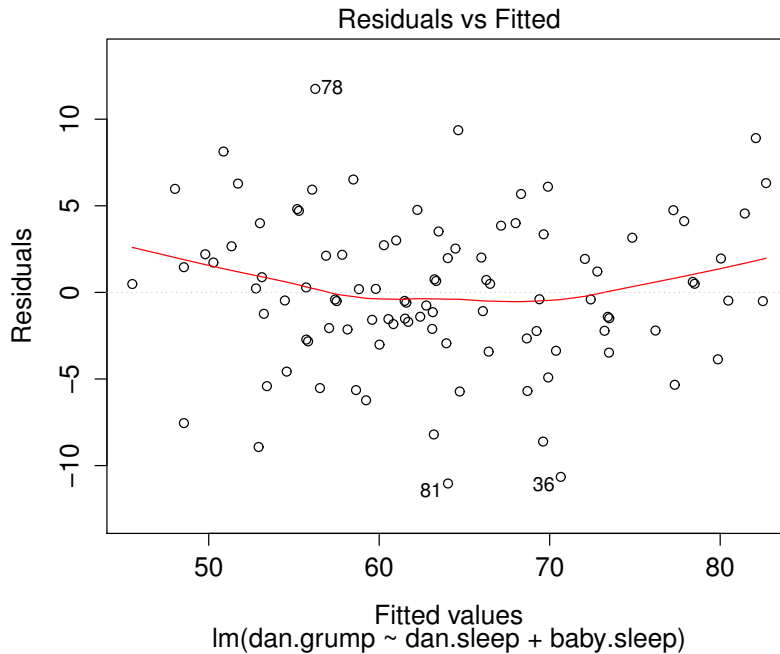


Figure 15.13: Plot of the fitted values against the residuals for `regression.2`, with a line showing the relationship between the two. If this is horizontal and straight, then we can feel reasonably confident that the “average residual” for all “fitted values” is more or less the same. This is one of the standard regression plots produced by the `plot()` function when the input is a linear regression object. It is obtained by setting `which=1`.

.....

and 15.14 is genuine;¹¹ although it still bears remembering that the pattern in Figure 15.12 is pretty damn straight: in other words the deviations from linearity are pretty small, and probably not worth worrying about.

In a lot of cases, the solution to this problem (and many others) is to transform one or more of the variables. We discussed the basics of variable transformation in Sections 7.2 and 7.3, but I do want to make special note of one additional possibility that I didn’t mention earlier: the Box-Cox transform. The Box-Cox function is a fairly simple one, but it’s very widely used

$$f(x, \lambda) = \frac{x^\lambda - 1}{\lambda}$$

for all values of λ except $\lambda = 0$. When $\lambda = 0$ we just take the natural logarithm (i.e., $\ln(x)$). You can calculate it using the `boxCox()` function in the `car` package. Better yet, if what you’re trying to do is convert a data to normal, or as normal as possible, there’s the `powerTransform()` function in the `car` package that can estimate the best value of λ . Variable transformation is another topic that deserves a fairly detailed treatment, but (again) due to deadline constraints, it will have to wait until a future version of this book.

¹¹And, if you take the time to check the `residualPlots()` for `regression.1`, it’s pretty clear that this isn’t some wacky distortion being caused by the fact that `baby.sleep` is a useless predictor variable. It’s an actual nonlinearity in the relationship between `dan.sleep` and `dan.grump`.

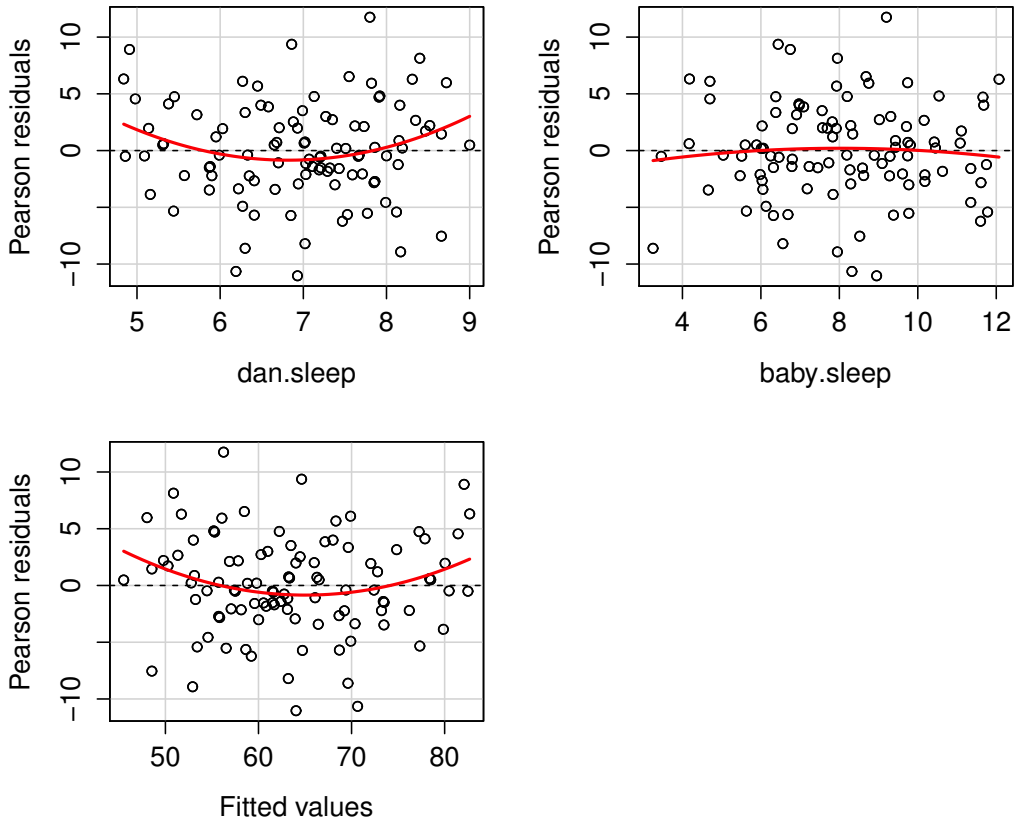


Figure 15.14: Plot of the fitted values against the residuals for `regression.2`, along with similar plots for the two predictors individually. This plot is produced by the `residualPlots()` function in the `car` package. Note that it refers to the residuals as “Pearson residuals”, but in this context these are the same as ordinary residuals.

15.8.5 Checking the homogeneity of variance

The regression models that we’ve talked about all make a homogeneity of variance assumption: the variance of the residuals is assumed to be constant. The “default” plot that R provides to help with doing this (`which = 3` when using `plot()`) shows a plot of the square root of the size of the residual $\sqrt{|\epsilon_i|}$, as a function of the fitted value \hat{Y}_i . We can produce the plot using the following command,

```
> plot(x = regression.2, which = 3)
```

and the resulting plot is shown in Figure 15.15. Note that this plot actually uses the standardised residuals (i.e., converted to z scores) rather than the raw ones, but it’s immaterial from our point of view. What we’re looking to see here is a straight, horizontal line running through the middle of the plot.

A slightly more formal approach is to run hypothesis tests. The `car` package provides a function called `ncvTest()` (**non-constant variance test**) that can be used for this purpose (Cook & Weisberg, 1983).

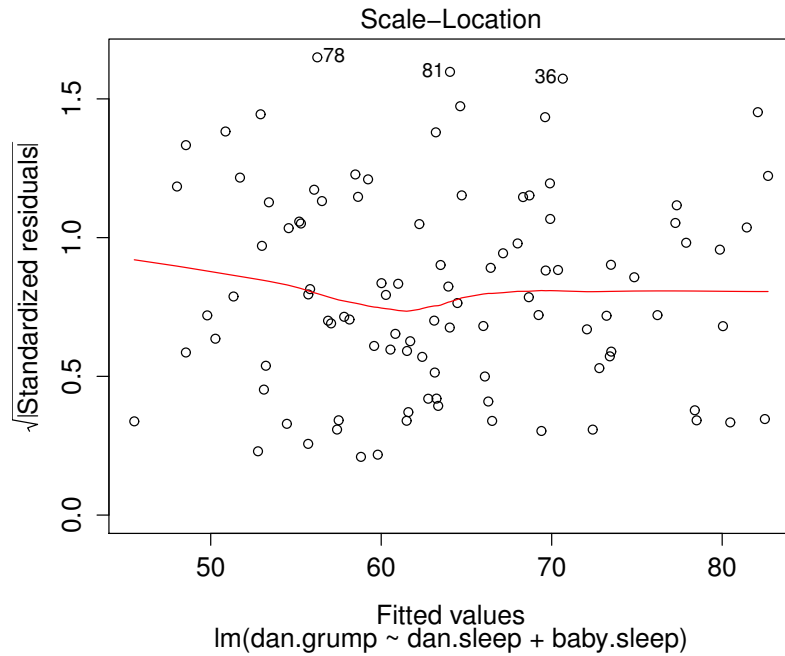


Figure 15.15: Plot of the fitted values (model predictions) against the square root of the abs standardised residuals. This plot is used to diagnose violations of homogeneity of variance. If the variance is really constant, then the line through the middle should be horizontal and flat. This is one of the standard regression plots produced by the `plot()` function when the input is a linear regression object. It is obtained by setting `which=3`.

.....

I won't explain the details of how it works, other than to say that the idea is that what you do is run a regression to see if there is a relationship between the squared residuals ϵ_i^2 and the fitted values \hat{Y}_i , or possibly to run a regression using all of the original predictors instead of just \hat{Y}_i .¹² Using the default settings, the `ncvTest()` looks for a relationship between \hat{Y}_i and the variance of the residuals, making it a straightforward analogue of Figure 15.15. So if we run it for our model,

```
> ncvTest( regression.2 )
Non-constant Variance Score Test
Variance formula: ~ fitted.values
Chisquare = 0.09317511    Df = 1      p = 0.7601788
```

We see that our original impression was right: there's no violations of homogeneity of variance in this data.

It's a bit beyond the scope of this chapter to talk too much about how to deal with violations of homogeneity of variance, but I'll give you a quick sense of what you need to consider. The *main* thing to worry about, if homogeneity of variance is violated, is that the standard error estimates associated with the regression coefficients are no longer entirely reliable, and so your *t* tests for the coefficients aren't

¹²Note that the underlying mechanics of the test aren't the same as the ones I've described for regressions; the goodness of fit is assessed using what's known as a score-test not an *F*-test, and the test statistic is (approximately) χ^2 distributed if there's no relationship

quite right either. A simple fix to the problem is to make use of a “heteroscedasticity corrected covariance matrix” when estimating the standard errors. These are often called **sandwich estimators**, for reasons that only make sense if you understand the maths at a low level¹³. You don’t need to understand what this means (not for an introductory class), but it might help to note that there’s a `hccm()` function in the `car()` package that does it. Better yet, you don’t even need to use it. You can use the `coeftest()` function in the `lmtest` package, but you need the `car` package loaded:

```
> coeftest( regression.2, vcov= hccm )

t test of coefficients:

            Estimate Std. Error t value Pr(>|t|)
(Intercept) 125.965566   3.247285 38.7910 <2e-16 ***
dan.sleep    -8.950250   0.615820 -14.5339 <2e-16 ***
baby.sleep     0.010524   0.291565   0.0361   0.9713
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

Not surprisingly, these t tests are pretty much identical to the ones that we saw when we used the `summary(regression.2)` command earlier; because the homogeneity of variance assumption wasn’t violated. But if it had been, we might have seen some more substantial differences.

15.8.6 Checking for collinearity

The last kind of regression diagnostic that I’m going to discuss in this chapter is the use of **variance inflation factors** (VIFs), which are useful for determining whether or not the predictors in your regression model are too highly correlated with each other. There is a variance inflation factor associated with each predictor X_k in the model, and the formula for the k -th VIF is:

$$\text{VIF}_k = \frac{1}{1 - R_{(-k)}^2}$$

where $R_{(-k)}^2$ refers to R -squared value you would get if you ran a regression using X_k as the outcome variable, and all the other X variables as the predictors. The idea here is that $R_{(-k)}^2$ is a very good measure of the extent to which X_k is correlated with all the other variables in the model. Better yet, the square root of the VIF is pretty interpretable: it tells you how much wider the confidence interval for the corresponding coefficient b_k is, relative to what you would have expected if the predictors are all nice and uncorrelated with one another. If you’ve only got two predictors, the VIF values are always going to be the same, as we can see if we use the `vif()` function (`car` package)...

```
> vif( mod = regression.2 )
dan.sleep baby.sleep
1.651038 1.651038
```

And since the square root of 1.65 is 1.28, we see that the correlation between our two predictors isn’t causing much of a problem.

¹³Again, a footnote that should be read only by the two readers of this book that love linear algebra (mmmm... I love the smell of matrix computations in the morning; smells like... nerd). In these estimators, the covariance matrix for \boldsymbol{b} is given by $(\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}'\boldsymbol{\Sigma}\mathbf{X} (\mathbf{X}'\mathbf{X})^{-1}$. See, it’s a “sandwich”? Assuming you think that $(\mathbf{X}'\mathbf{X})^{-1}$ = “bread” and $\mathbf{X}'\boldsymbol{\Sigma}\mathbf{X}$ = “filling”, that is. Which of course everyone does, right? In any case, the usual estimator is what you get when you set $\boldsymbol{\Sigma} = \hat{\sigma}^2 \mathbf{I}$. The corrected version that I learned originally uses $\boldsymbol{\Sigma} = \text{diag}(\epsilon_i^2)$ (White, 1980). However, the version that Fox and Weisberg (2011) have implemented as the default in the `hccm()` function is a tweak on this, proposed by Long and Ervin (2000). This version uses $\boldsymbol{\Sigma} = \text{diag}(\epsilon_i^2 / (1 - h_i^2))$, where h_i is the i th hat value. Gosh, regression is *fun*, isn’t it?

To give a sense of how we could end up with a model that has bigger collinearity problems, suppose I were to run a much less interesting regression model, in which I tried to predict the `day` on which the data were collected, as a function of all the other variables in the data set. To see why this would be a bit of a problem, let's have a look at the correlation matrix for all four variables:

```
> cor( parenthood )
      dan.sleep  baby.sleep  dan.grump       day
dan.sleep    1.0000000  0.62794934 -0.90338404 -0.09840768
baby.sleep   0.62794934  1.00000000 -0.56596373 -0.01043394
dan.grump   -0.90338404 -0.56596373  1.00000000  0.07647926
day        -0.09840768 -0.01043394  0.07647926  1.00000000
```

We have some fairly large correlations between some of our predictor variables! When we run the regression model and look at the VIF values, we see that the collinearity is causing a lot of uncertainty about the coefficients. First, run the regression...

```
> regression.3 <- lm( day ~ baby.sleep + dan.sleep + dan.grump, parenthood )
```

and second, look at the VIFs...

```
> vif( regression.3 )
baby.sleep  dan.sleep  dan.grump
1.651064    6.102337  5.437903
```

Yep, that's some mighty fine collinearity you've got there.

15.9

Model selection

One fairly major problem that remains is the problem of “model selection”. That is, if we have a data set that contains several variables, which ones should we include as predictors, and which ones should we not include? In other words, we have a problem of **variable selection**. In general, model selection is a complex business, but it's made somewhat simpler if we restrict ourselves to the problem of choosing a subset of the variables that ought to be included in the model. Nevertheless, I'm not going to try covering even this reduced topic in a lot of detail. Instead, I'll talk about two broad principles that you need to think about; and then discuss one concrete tool that R provides to help you select a subset of variables to include in your model. Firstly, the two principles:

- It's nice to have an actual substantive basis for your choices. That is, in a lot of situations you the researcher have good reasons to pick out a smallish number of possible regression models that are of theoretical interest; these models will have a sensible interpretation in the context of your field. Never discount the importance of this. Statistics serves the scientific process, not the other way around.
- To the extent that your choices rely on statistical inference, there is a trade off between simplicity and goodness of fit. As you add more predictors to the model, you make it more complex; each predictor adds a new free parameter (i.e., a new regression coefficient), and each new parameter increases the model's capacity to “absorb” random variations. So the goodness of fit (e.g., R^2) continues to rise as you add more predictors no matter what. If you want your model to be able to generalise well to new observations, you need to avoid throwing in too many variables.

This latter principle is often referred to as **Ockham's razor**, and is often summarised in terms of the following pithy saying: *do not multiply entities beyond necessity*. In this context, it means: don't chuck in a bunch of largely irrelevant predictors just to boost your R^2 . Hm. Yeah, the original was better.

In any case, what we need is an actual mathematical criterion that will implement the qualitative principle behind Ockham's razor in the context of selecting a regression model. As it turns out there are several possibilities. The one that I'll talk about is the **Akaike information criterion** (AIC; Akaike, 1974) simply because it's the default one used in the R function `step()`. In the context of a linear regression model (and ignoring terms that don't depend on the model in any way!), the AIC for a model that has K predictor variables plus an intercept is:¹⁴

$$\text{AIC} = \frac{\text{SS}_{\text{res}}}{\hat{\sigma}^2} + 2K$$

The smaller the AIC value, the better the model performance is. If we ignore the low level details, it's fairly obvious what the AIC does: on the left we have a term that increases as the model predictions get worse; on the right we have a term that increases as the model complexity increases. The best model is the one that fits the data well (low residuals; left hand side) using as few predictors as possible (low K ; right hand side). In short, this is a simple implementation of Ockham's razor.

15.9.1 Backward elimination

Okay, let's have a look at the `step()` function at work. In this example I'll keep it simple and use only the basic **backward elimination** approach. That is, start with the complete regression model, including all possible predictors. Then, at each "step" we try all possible ways of removing one of the variables, and whichever of these is best (in terms of lowest AIC value) is accepted. This becomes our new regression model; and we then try all possible deletions from the new model, again choosing the option with lowest AIC. This process continues until we end up with a model that has a lower AIC value than any of the other possible models that you could produce by deleting one of its predictors. Let's see this in action. First, I need to define the model from which the process starts.

```
> full.model <- lm( formula = dan.grump ~ dan.sleep + baby.sleep + day,
+                     data = parenthood
+ )
```

That's nothing terribly new: yet another regression. Booooring. Still, we do need to do it: the `object` argument to the `step()` function will be this regression model. With this in mind, I would call the `step()` function using the following command:

```
> step( object = full.model,      # start at the full model
+        direction = "backward"   # allow it remove predictors but not add them
+ )
```

although in practice I didn't need to specify `direction` because "`backward`" is the default. The output is somewhat lengthy, so I'll go through it slowly. Firstly, the output reports the AIC value for the current best model:

```
Start: AIC=299.08
dan.grump ~ dan.sleep + baby.sleep + day
```

¹⁴Note, however, that the `step()` function computes the full version of AIC, including the irrelevant constants that I've dropped here. As a consequence this equation won't correctly describe the AIC values that you see in the outputs here. However, if you calculate the AIC values using my formula for two different regression models and take the difference between them, this will be the same as the differences between AIC values that `step()` reports. In practice, this is all you care about: the actual value of an AIC statistic isn't very informative, but the differences between two AIC values *are* useful, since these provide a measure of the extent to which one model outperforms another.

That's our starting point. Since small AIC values are good, we want to see if we can get a value smaller than 299.08 by deleting one of those three predictors. So what R does is try all three possibilities, calculate the AIC values for each one, and then print out a short table with the results:

	Df	Sum of Sq	RSS	AIC
- baby.sleep	1	0.1	1837.2	297.08
- day	1	1.6	1838.7	297.16
<none>			1837.1	299.08
- dan.sleep	1	4909.0	6746.1	427.15

To read this table, it helps to note that the text in the left hand column is telling you what *change* R made to the regression model. So the line that reads <none> is the actual model we started with, and you can see on the right hand side that this still corresponds to an AIC value of 299.08 (obviously). The other three rows in the table correspond to the other three models that it looked at: it tried removing the `baby.sleep` variable, which is indicated by - `baby.sleep`, and this produced an AIC value of 297.08. That was the best of the three moves, so it's at the top of the table. So, this move is accepted, and now we start again. There are two predictors left in the model, `dan.sleep` and `day`, so it tries deleting those:

```
Step: AIC=297.08
dan.grump ~ dan.sleep + day
```

	Df	Sum of Sq	RSS	AIC
- day	1	1.6	1838.7	295.17
<none>			1837.2	297.08
- dan.sleep	1	8103.0	9940.1	463.92

Okay, so what we can see is that removing the `day` variable lowers the AIC value from 297.08 to 295.17. So R decides to keep that change too, and moves on:

```
Step: AIC=295.17
dan.grump ~ dan.sleep
```

	Df	Sum of Sq	RSS	AIC
<none>		1838.7	295.17	
- dan.sleep	1	8159.9	9998.6	462.50

This time around, there's no further deletions that can actually improve the AIC value. So the `step()` function stops, and prints out the result of the best regression model it could find:

```
Call:
lm(formula = dan.grump ~ dan.sleep, data = parenthood)

Coefficients:
(Intercept)    dan.sleep
      125.956       -8.937
```

which is (perhaps not all that surprisingly) the `regression.1` model that we started with at the beginning of the chapter.

15.9.2 Forward selection

As an alternative, you can also try **forward selection**. This time around we start with the smallest possible model as our start point, and only consider the possible additions to the model. However, there's

one complication: you also need to tell `step()` what the largest possible model you're willing to entertain is, using the `scope` argument. The simplest usage is like this:

```
> null.model <- lm( dan.grump ~ 1, parenthood ) # intercept only.
> step( object = null.model,          # start with null.model
+       direction = "forward",      # only consider "addition" moves
+       scope = dan.grump ~ dan.sleep + baby.sleep + day # largest model allowed
+ )
```

If I do this, the output takes on a similar form, but now it only considers addition (+) moves rather than deletion (-) moves:

```
Start: AIC=462.5
dan.grump ~ 1

      Df Sum of Sq   RSS   AIC
+ dan.sleep    1     8159.9 1838.7 295.17
+ baby.sleep   1     3202.7 6795.9 425.89
<none>                   9998.6 462.50
+ day          1      58.5 9940.1 463.92

Step: AIC=295.17
dan.grump ~ dan.sleep

      Df Sum of Sq   RSS   AIC
<none>                   1838.7 295.17
+ day          1    1.55760 1837.2 297.08
+ baby.sleep   1    0.02858 1838.7 297.16

Call:
lm(formula = dan.grump ~ dan.sleep, data = parenthood)

Coefficients:
(Intercept)  dan.sleep
        125.956      -8.937
```

As you can see, it's found the same model. In general though, forward and backward selection don't always have to end up in the same place.

15.9.3 A **caveat**

Automated variable selection methods are seductive things, especially when they're bundled up in (fairly) simple functions like `step()`. They provide an element of objectivity to your model selection, and that's kind of nice. Unfortunately, they're sometimes used as an excuse for thoughtlessness. No longer do you have to think carefully about which predictors to add to the model and what the theoretical basis for their inclusion might be... everything is solved by the magic of AIC. And if we start throwing around phrases like Ockham's razor, well, it sounds like everything is wrapped up in a nice neat little package that no-one can argue with.

Or, perhaps not. Firstly, there's very little agreement on what counts as an appropriate model selection criterion. When I was taught backward elimination as an undergraduate, we used *F*-tests to do it, because that was the default method used by the software. The default in the `step()` function is AIC, and since this is an introductory text that's the only method I've described, but the AIC is hardly the Word of the Gods of Statistics. It's an approximation, derived under certain assumptions, and it's

guaranteed to work only for large samples when those assumptions are met. Alter those assumptions and you get a different criterion, like the BIC for instance. Take a different approach again and you get the NML criterion. Decide that you're a Bayesian and you get model selection based on posterior odds ratios. Then there are a bunch of regression specific tools that I haven't mentioned. And so on. All of these different methods have strengths and weaknesses, and some are easier to calculate than others (AIC is probably the easiest of the lot, which might account for its popularity). Almost all of them produce the same answers when the answer is "obvious" but there's a fair amount of disagreement when the model selection problem becomes hard.

What does this mean in practice? Well, you *could* go and spend several years teaching yourself the theory of model selection, learning all the ins and outs of it; so that you could finally decide on what you personally think the right thing to do is. Speaking as someone who actually did that, I wouldn't recommend it: you'll probably come out the other side even more confused than when you started. A better strategy is to show a bit of common sense... if you're staring at the results of a `step()` procedure, and the model that makes sense is close to having the smallest AIC, but is narrowly defeated by a model that doesn't make any sense... trust your instincts. Statistical model selection is an inexact tool, and as I said at the beginning, *interpretability matters*.

15.9.4 Comparing two regression models

An alternative to using automated model selection procedures is for the researcher to explicitly select two or more regression models to compare to each other. You can do this in a few different ways, depending on what research question you're trying to answer. Suppose we want to know whether or not the amount of sleep that my son got has any relationship to my grumpiness, over and above what we might expect from the amount of sleep that I got. We also want to make sure that the day on which we took the measurement has no influence on the relationship. That is, we're interested in the relationship between `baby.sleep` and `dan.grump`, and from that perspective `dan.sleep` and `day` are nuisance variable or **covariates** that we want to control for. In this situation, what we would like to know is whether `dan.grump ~ dan.sleep + day + baby.sleep` (which I'll call Model 1, or `M1`) is a better regression model for these data than `dan.grump ~ dan.sleep + day` (which I'll call Model 0, or `M0`). There are two different ways we can compare these two models, one based on a model selection criterion like AIC, and the other based on an explicit hypothesis test. I'll show you the AIC based approach first because it's simpler, and follows naturally from the `step()` function that we saw in the last section. The first thing I need to do is actually run the regressions:

```
> M0 <- lm( dan.grump ~ dan.sleep + day, parenthood )
> M1 <- lm( dan.grump ~ dan.sleep + day + baby.sleep, parenthood )
```

Now that I have my regression models, I could use the `summary()` function to run various hypothesis tests and other useful statistics, just as we have discussed throughout this chapter. However, since the current focus on model comparison, I'll skip this step and go straight to the AIC calculations. Conveniently, the `AIC()` function in R lets you input several regression models, and it will spit out the AIC values for each of them.¹⁵

```
> AIC( M0, M1 )
      df      AIC
M0  4  582.8681
M1  5  584.8646
```

¹⁵While I'm on this topic I should point out that there is also a function called `BIC()` which computes the Bayesian information criterion (BIC) for the models. So you could type `BIC(M0,M1)` and get a very similar output. In fact, while I'm not particularly impressed with either AIC or BIC as model selection methods, if you do find yourself using one of these two, the empirical evidence suggests that BIC is the better criterion of the two. In most simulation studies that I've seen, BIC does a much better job of selecting the correct model.

Since Model 0 has the smaller AIC value, it is judged to be the better model for these data.

A somewhat different approach to the problem comes out of the hypothesis testing framework. Suppose you have two regression models, where one of them (Model 0) contains a *subset* of the predictors from the other one (Model 1). That is, Model 1 contains all of the predictors included in Model 0, plus one or more additional predictors. When this happens we say that Model 0 is **nested** within Model 1, or possibly that Model 0 is a **submodel** of Model 1. Regardless of the terminology what this means is that we can think of Model 0 as a null hypothesis and Model 1 as an alternative hypothesis. And in fact we can construct an F test for this in a fairly straightforward fashion. We can fit both models to the data and obtain a residual sum of squares for both models. I'll denote these as $SS_{res}^{(0)}$ and $SS_{res}^{(1)}$ respectively. The superscripting here just indicates which model we're talking about. Then our F statistic is

$$F = \frac{(SS_{res}^{(0)} - SS_{res}^{(1)})/k}{(SS_{res}^{(1)})/(N - p - 1)}$$

where N is the number of observations, p is the number of predictors in the full model (not including the intercept), and k is the difference in the number of parameters between the two models.¹⁶ The degrees of freedom here are k and $N - p - 1$. Note that it's often more convenient to think about the difference between those two SS values as a sum of squares in its own right. That is:

$$SS_{\Delta} = SS_{res}^{(0)} - SS_{res}^{(1)}$$

The reason why this is helpful is that we can express SS_{Δ} a measure of the extent to which the two models make different predictions about the outcome variable. Specifically:

$$SS_{\Delta} = \sum_i (\hat{y}_i^{(1)} - \hat{y}_i^{(0)})^2$$

where $\hat{y}_i^{(0)}$ is the fitted value for y_i according to model M_0 and $\hat{y}_i^{(1)}$ is the fitted value for y_i according to model M_1 .

Okay, so that's the hypothesis test that we use to compare two regression models to one another. Now, how do we do it in R? The answer is to use the `anova()` function. All we have to do is input the two models that we want to compare (null model first):

```
> anova( M0, M1 )
Analysis of Variance Table

Model 1: dan.grump ~ dan.sleep + day
Model 2: dan.grump ~ dan.sleep + day + baby.sleep
  Res.Df   RSS Df Sum of Sq    F Pr(>F)
1     97 1837.2
2     96 1837.1  1  0.063688 0.0033 0.9541
```

Note that, just like we saw with the output from the `step()` function, R has used the acronym RSS to refer to the residual sum of squares from each model. That is, RSS in this output corresponds to SS_{res} in the formula above. Since we have $p > .05$ we retain the null hypothesis (**M0**). This approach to regression, in which we add all of our covariates into a null model, and then *add* the variables of interest into an alternative model, and then compare the two models in hypothesis testing framework, is often referred to as **hierarchical regression**.

¹⁶It's worth noting in passing that this same F statistic can be used to test a much broader range of hypotheses than those that I'm mentioning here. Very briefly: notice that the nested model M_0 corresponds to the full model M_1 when we constrain some of the regression coefficients to zero. It is sometimes useful to construct submodels by placing other kinds of constraints on the regression coefficients. For instance, maybe two different coefficients might have to sum to zero, or something like that. You can construct hypothesis tests for those kind of constraints too, but it is somewhat more complicated and the sampling distribution for F can end up being something known as the non-central F distribution, which is waaaaay beyond the scope of this book! All I want to do is alert you to this possibility.

Summary

- Basic ideas in linear regression and how regression models are estimated (Sections 15.1 and 15.2).
- Multiple linear regression (Section 15.3).
- Measuring the overall performance of a regression model using R^2 (Section 15.4)
- Hypothesis tests for regression models (Section 15.5)
- Calculating confidence intervals for regression coefficients, and standardised coefficients (Section 15.6)
- The assumptions of regression (Section 15.7) and how to check them (Section 15.8)
- Selecting a regression model (Section 15.9)

16. Factorial ANOVA

Over the course of the last few chapters you can probably detect a general trend. We started out looking at tools that you can use to compare two groups to one another, most notably the *t*-test (Chapter 13). Then, we introduced analysis of variance (ANOVA) as a method for comparing more than two groups (Chapter 14). The chapter on regression (Chapter 15) covered a somewhat different topic, but in doing so it introduced a powerful new idea: building statistical models that have *multiple* predictor variables used to explain a single outcome variable. For instance, a regression model could be used to predict the number of errors a student makes in a reading comprehension test based on the number of hours they studied for the test, and their score on a standardised IQ test. The goal in this chapter is to import this idea into the ANOVA framework. For instance, suppose we were interested in using the reading comprehension test to measure student achievements in three different schools, and we suspect that girls and boys are developing at different rates (and so would be expected to have different performance on average). Each student is classified in two different ways: on the basis of their gender, and on the basis of their school. What we'd like to do is analyse the reading comprehension scores in terms of *both* of these grouping variables. The tool for doing so is generically referred to as **factorial ANOVA**. However, since we have two grouping variables, we sometimes refer to the analysis as a two-way ANOVA, in contrast to the one-way ANOVAs that we ran in Chapter 14.

16.1

Factorial ANOVA 1: balanced designs, no interactions

When we discussed analysis of variance in Chapter 14, we assumed a fairly simple experimental design: each person falls into one of several groups, and we want to know whether these groups have different means on some outcome variable. In this section, I'll discuss a broader class of experimental designs, known as **factorial designs**, in which we have more than one grouping variable. I gave one example of how this kind of design might arise above. Another example appears in Chapter 14, in which we were looking at the effect of different drugs on the `mood.gain` experienced by each person. In that chapter we did find a significant effect of drug, but at the end of the chapter we also ran an analysis to see if there was an effect of therapy. We didn't find one, but there's something a bit worrying about trying to run two *separate* analyses trying to predict the same outcome. Maybe there actually *is* an effect of therapy on mood gain, but we couldn't find it because it was being "hidden" by the effect of drug? In other words, we're going to want to run a *single* analysis that includes *both* `drug` and `therapy` as predictors. For this analysis each person is cross-classified by the drug they were given (a factor with 3 levels) and what therapy they received (a factor with 2 levels). We refer to this as a 3×2 factorial design. If we cross-tabulate `drug` by `therapy`, using the `xtabs()` function (see Section 7.1), we get the following table:

```
> xtabs(~ drug + therapy, clin.trial)
      therapy
drug      no.therapy CBT
placebo      3   3
anxifree     3   3
joyzepam     3   3
```

As you can see, not only do we have participants corresponding to all possible combinations of the two factors, indicating that our design is **completely crossed**, it turns out that there are an equal number of people in each group. In other words, we have a **balanced** design. In this section I'll talk about how to analyse data from balanced designs, since this is the simplest case. The story for unbalanced designs is quite tedious, so we'll put it to one side for the moment.

16.1.1 What hypotheses are we testing?

Like one-way ANOVA, factorial ANOVA is a tool for testing certain types of hypotheses about population means. So a sensible place to start would be to be explicit about what our hypotheses actually are. However, before we can even get to that point, it's really useful to have some clean and simple notation to describe the population means. Because of the fact that observations are cross-classified in terms of two different factors, there are quite a lot of different means that one might be interested. To see this, let's start by thinking about all the different sample means that we can calculate for this kind of design. Firstly, there's the obvious idea that we might be interested in this table of group means:

```
> aggregate(mood.gain ~ drug + therapy, clin.trial, mean)
      drug      therapy mood.gain
1 placebo no.therapy  0.300000
2 anxifree no.therapy  0.400000
3 joyzepam no.therapy  1.466667
4 placebo      CBT  0.600000
5 anxifree      CBT  1.033333
6 joyzepam      CBT  1.500000
```

Now, this output shows a cross-tabulation of the group means for all possible combinations of the two factors (e.g., people who received the placebo and no therapy, people who received the placebo while getting CBT, etc). However, we can also construct tables that ignore one of the two factors. That gives us output that looks like this:

```
> aggregate(mood.gain ~ drug, clin.trial, mean)
      drug mood.gain
1 placebo 0.4500000
2 anxifree 0.7166667
3 joyzepam 1.4833333

> aggregate(mood.gain ~ therapy, clin.trial, mean)
      therapy mood.gain
1 no.therapy 0.7222222
2      CBT  1.0444444
```

But of course, if we can ignore one factor we can certainly ignore both. That is, we might also be interested in calculating the average mood gain across all 18 participants, regardless of what drug or psychological therapy they were given:

```
> mean(clin.trial$mood.gain)
[1] 0.88333
```

At this point we have 12 different sample means to keep track of! It is helpful to organise all these numbers into a single table, which would look like this:

	no therapy	CBT	total
placebo	0.30	0.60	0.45
anxitfree	0.40	1.03	0.72
joyzepam	1.47	1.50	1.48
total	0.72	1.04	0.88

Now, each of these different means is of course a sample statistic: it's a quantity that pertains to the specific observations that we've made during our study. What we want to make inferences about are the corresponding population parameters: that is, the true means as they exist within some broader population. Those population means can also be organised into a similar table, but we'll need a little mathematical notation to do so. As usual, I'll use the symbol μ to denote a population mean. However, because there are lots of different means, I'll need to use subscripts to distinguish between them.

Here's how the notation works. Our table is defined in terms of two factors: each row corresponds to a different level of Factor A (in this case `drug`), and each column corresponds to a different level of Factor B (in this case `therapy`). If we let R denote the number of rows in the table, and C denote the number of columns, we can refer to this as an $R \times C$ factorial ANOVA. In this case $R = 3$ and $C = 2$. We'll use lowercase letters to refer to specific rows and columns, so μ_{rc} refers to the population mean associated with the r th level of Factor A (i.e. row number r) and the c th level of Factor B (column number c).¹ So the population means are now written like this:

	no therapy	CBT	total
placebo	μ_{11}	μ_{12}	
anxitfree	μ_{21}	μ_{22}	
joyzepam	μ_{31}	μ_{32}	
total			

Okay, what about the remaining entries? For instance, how should we describe the average mood gain across the entire (hypothetical) population of people who might be given Joyzepam in an experiment like this, regardless of whether they were in CBT? We use the “dot” notation to express this. In the case of Joyzepam, notice that we're talking about the mean associated with the third row in the table. That is, we're averaging across two cell means (i.e., μ_{31} and μ_{32}). The result of this averaging is referred to as a **marginal mean**, and would be denoted μ_3 in this case. The marginal mean for CBT corresponds to the population mean associated with the second column in the table, so we use the notation $\mu_{.2}$ to describe it. The grand mean is denoted $\mu_{..}$ because it is the mean obtained by averaging (marginalising²) over both. So our full table of population means can be written down like this:

	no therapy	CBT	total
placebo	μ_{11}	μ_{12}	$\mu_{1.}$
anxitfree	μ_{21}	μ_{22}	$\mu_{2.}$
joyzepam	μ_{31}	μ_{32}	$\mu_{3.}$
total	$\mu_{.1}$	$\mu_{.2}$	$\mu_{..}$

¹The nice thing about the subscript notation is that generalises nicely: if our experiment had involved a third factor, then we could just add a third subscript. In principle, the notation extends to as many factors as you might care to include, but in this book we'll rarely consider analyses involving more than two factors, and never more than three.

²Technically, marginalising isn't quite identical to a regular mean: it's a weighted average, where you take into account the frequency of the different events that you're averaging over. However, in a balanced design, all of our cell frequencies are equal by definition, so the two are equivalent. We'll discuss unbalanced designs later, and when we do so you'll see that all of our calculations become a real headache. But let's ignore this for now.

Now that we have this notation, it is straightforward to formulate and express some hypotheses. Let's suppose that the goal is to find out two things: firstly, does the choice of drug have any effect on mood, and secondly, does CBT have any effect on mood? These aren't the only hypotheses that we could formulate of course, and we'll see a really important example of a different kind of hypothesis in Section 16.2, but these are the two simplest hypotheses to test, and so we'll start there. Consider the first test. If drug has no effect, then we would expect all of the row means to be identical, right? So that's our null hypothesis. On the other hand, if the drug does matter then we should expect these row means to be different. Formally, we write down our null and alternative hypotheses in terms of the *equality of marginal means*:

Null hypothesis, H_0 : row means are the same, i.e., $\mu_1 = \mu_2 = \mu_3$.
Alternative hypothesis, H_1 : at least one row mean is different.

It's worth noting that these are *exactly* the same statistical hypotheses that we formed when we ran a one-way ANOVA on these data back in Chapter 14. Back then I used the notation μ_P to refer to the mean mood gain for the placebo group, with μ_A and μ_J corresponding to the group means for the two drugs, and the null hypothesis was $\mu_P = \mu_A = \mu_J$. So we're actually talking about the same hypothesis: it's just that the more complicated ANOVA requires more careful notation due to the presence of multiple grouping variables, so we're now referring to this hypothesis as $\mu_1 = \mu_2 = \mu_3$. However, as we'll see shortly, although the hypothesis is identical, the test of that hypothesis is subtly different due to the fact that we're now acknowledging the existence of the second grouping variable.

Speaking of the other grouping variable, you won't be surprised to discover that our second hypothesis test is formulated the same way. However, since we're talking about the psychological therapy rather than drugs, our null hypothesis now corresponds to the equality of the column means:

Null hypothesis, H_0 : column means are the same, i.e., $\mu_{.1} = \mu_{.2}$
Alternative hypothesis, H_1 : column means are different, i.e., $\mu_{.1} \neq \mu_{.2}$

16.1.2 Running the analysis in R

The null and alternative hypotheses that I described in the last section should seem awfully familiar: they're basically the same as the hypotheses that we were testing in our simpler one-way ANOVAs in Chapter 14. So you're probably expecting that the hypothesis *tests* that are used in factorial ANOVA will be essentially the same as the *F*-test from Chapter 14. You're expecting to see references to sums of squares (SS), mean squares (MS), degrees of freedom (df), and finally an *F*-statistic that we can convert into a *p*-value, right? Well, you're absolutely and completely right. So much so that I'm going to depart from my usual approach. Throughout this book, I've generally taken the approach of describing the logic (and to an extent the mathematics) that underpins a particular analysis first; and only then introducing the R commands that you'd use to produce the analysis. This time I'm going to do it the other way around, and show you the R commands first. The reason for doing this is that I want to highlight the similarities between the simple one-way ANOVA tool that we discussed in Chapter 14, and the more complicated tools that we're going to use in this chapter.

If the data you're trying to analyse correspond to a balanced factorial design, then running your analysis of variance is easy. To see how easy it is, let's start by reproducing the original analysis from Chapter 14. In case you've forgotten, for that analysis we were using only a single factor (i.e., `drug`) to predict our outcome variable (i.e., `mood.gain`), and so this was what we did:

```
> model.1 <- aov( mood.gain ~ drug, clin.trial )
> summary( model.1 )
   Df Sum Sq Mean Sq F value    Pr(>F)
```

```

drug      2   3.45   1.727    18.6 8.6e-05 ***
Residuals 15   1.39   0.093
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1

```

Note that this time around I've used the name `model.1` as the label for my `aov` object, since I'm planning on creating quite a few other models too. To start with, suppose I'm also curious to find out if `therapy` has a relationship to `mood.gain`. In light of what we've seen from our discussion of multiple regression in Chapter 15, you probably won't be surprised that all we have to do is extend the formula: in other words, if we specify `mood.gain ~ drug + therapy` as our model, we'll probably get what we're after:

```

> model.2 <- aov( mood.gain ~ drug + therapy, clin.trial )
> summary( model.2 )
   Df Sum Sq Mean Sq F value Pr(>F)
drug      2   3.45   1.727    26.15 1.9e-05 ***
therapy    1   0.47   0.467     7.08   0.019 *
Residuals 14   0.92   0.066
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1

```

This output is pretty simple to read too: the first row of the table reports a between-group sum of squares (SS) value associated with the `drug` factor, along with a corresponding between-group *df* value. It also calculates a mean square value (MS), and *F*-statistic and a *p*-value. There is also a row corresponding to the `therapy` factor, and a row corresponding to the residuals (i.e., the within groups variation).

Not only are all of the individual quantities pretty familiar, the relationships between these different quantities has remained unchanged: just like we saw with the original one-way ANOVA, note that the mean square value is calculated by dividing SS by the corresponding *df*. That is, it's still true that

$$MS = \frac{SS}{df}$$

regardless of whether we're talking about `drug`, `therapy` or the residuals. To see this, let's not worry about how the sums of squares values are calculated: instead, let's take it on faith that R has calculated the SS values correctly, and try to verify that all the rest of the numbers make sense. First, note that for the `drug` factor, we divide 3.45 by 2, and end up with a mean square value of 1.73. For the `therapy` factor, there's only 1 degree of freedom, so our calculations are even simpler: dividing 0.47 (the SS value) by 1 gives us an answer of 0.47 (the MS value).

Turning to the *F* statistics and the *p* values, notice that we have two of each: one corresponding to the `drug` factor and the other corresponding to the `therapy` factor. Regardless of which one we're talking about, the *F* statistic is calculated by dividing the mean square value associated with the factor by the mean square value associated with the residuals. If we use "A" as shorthand notation to refer to the first factor (factor A; in this case `drug`) and "R" as shorthand notation to refer to the residuals, then the *F* statistic associated with factor A is denoted F_A , and is calculated as follows:

$$F_A = \frac{MS_A}{MS_R}$$

and an equivalent formula exists for factor B (i.e., `therapy`). Note that this use of "R" to refer to residuals is a bit awkward, since we also used the letter R to refer to the number of rows in the table, but I'm only going to use "R" to mean residuals in the context of SS_R and MS_R , so hopefully this shouldn't be confusing. Anyway, to apply this formula to the `drugs` factor, we take the mean square of 1.73 and divide it by the residual mean square value of 0.07, which gives us an *F*-statistic of 26.15. The corresponding calculation for the `therapy` variable would be to divide 0.47 by 0.07 which gives 7.08 as the *F*-statistic. Not surprisingly, of course, these are the same values that R has reported in the ANOVA table above.

The last part of the ANOVA table is the calculation of the p values. Once again, there is nothing new here: for each of our two factors, what we're trying to do is test the null hypothesis that there is no relationship between the factor and the outcome variable (I'll be a bit more precise about this later on). To that end, we've (apparently) followed a similar strategy that we did in the one way ANOVA, and have calculated an F -statistic for each of these hypotheses. To convert these to p values, all we need to do is note that the sampling distribution for the F statistic under the null hypothesis (that the factor in question is irrelevant) is an F distribution: and that two degrees of freedom values are those corresponding to the factor, and those corresponding to the residuals. For the `drug` factor we're talking about an F distribution with 2 and 14 degrees of freedom (I'll discuss degrees of freedom in more detail later). In contrast, for the `therapy` factor sampling distribution is F with 1 and 14 degrees of freedom. If we really wanted to, we could calculate the p value ourselves using the `pf()` function (see Section 9.5). Just to prove that there's nothing funny going on, here's what that would look like for the `drug` variable:

```
> pf( q=26.15, df1=2, df2=14, lower.tail=FALSE )
[1] 1.871981e-05
```

And as you can see, this is indeed the p value reported in the ANOVA table above.

At this point, I hope you can see that the ANOVA table for this more complicated analysis corresponding to `model.2` should be read in much the same way as the ANOVA table for the simpler analysis for `model.1`. In short, it's telling us that the factorial ANOVA for our 3×2 design found a significant effect of drug ($F_{2,14} = 26.15, p < .001$) as well as a significant effect of therapy ($F_{1,14} = 7.08, p = .02$). Or, to use the more technically correct terminology, we would say that there are two **main effects** of drug and therapy. At the moment, it probably seems a bit redundant to refer to these as "main" effects: but it actually does make sense. Later on, we're going to want to talk about the possibility of "interactions" between the two factors, and so we generally make a distinction between main effects and interaction effects.

16.1.3 How are the sum of squares calculated?

In the previous section I had two goals: firstly, to show you that the R commands needed to do factorial ANOVA are pretty much the same ones that we used for a one way ANOVA. The only difference is the `formula` argument to the `aov()` function. Secondly, I wanted to show you what the ANOVA table looks like in this case, so that you can see from the outset that the basic logic and structure behind factorial ANOVA is the same as that which underpins one way ANOVA. Try to hold onto that feeling. It's genuinely true, insofar as factorial ANOVA is built in more or less the same way as the simpler one-way ANOVA model. It's just that this feeling of familiarity starts to evaporate once you start digging into the details. Traditionally, this comforting sensation is replaced by an urge to murder the authors of statistics textbooks.

Okay, let's start looking at some of those details. The explanation that I gave in the last section illustrates the fact that the hypothesis tests for the main effects (of drug and therapy in this case) are F -tests, but what it doesn't do is show you how the sum of squares (SS) values are calculated. Nor does it tell you explicitly how to calculate degrees of freedom (df values) though that's a simple thing by comparison. Let's assume for now that we have only two predictor variables, Factor A and Factor B. If we use Y to refer to the outcome variable, then we would use Y_{rci} to refer to the outcome associated with the i -th member of group rc (i.e., level/row r for Factor A and level/column c for Factor B). Thus, if we use \bar{Y} to refer to a sample mean, we can use the same notation as before to refer to group means, marginal means and grand means: that is, \bar{Y}_{rc} is the sample mean associated with the r th level of Factor A and the c th level of Factor B, $\bar{Y}_{r..}$ would be the marginal mean for the r th level of Factor A, $\bar{Y}_{..c}$ would be the marginal mean for the c th level of Factor B, and $\bar{Y}_{...}$ is the grand mean. In other words, our sample

means can be organised into the same table as the population means. For our clinical trial data, that table looks like this:

	no therapy	CBT	total
placebo	\bar{Y}_{11}	\bar{Y}_{12}	$\bar{Y}_{1..}$
anxitfree	\bar{Y}_{21}	\bar{Y}_{22}	$\bar{Y}_{2..}$
joyzepam	\bar{Y}_{31}	\bar{Y}_{32}	$\bar{Y}_{3..}$
total	$\bar{Y}_{..1}$	$\bar{Y}_{..2}$	$\bar{Y}_{..}$

And if we look at the sample means that I showed earlier, we have $\bar{Y}_{11} = 0.30$, $\bar{Y}_{12} = 0.60$ etc. In our clinical trial example, the `drugs` factor has 3 levels and the `therapy` factor has 2 levels, and so what we're trying to run is a 3×2 factorial ANOVA. However, we'll be a little more general and say that Factor A (the row factor) has R levels and Factor B (the column factor) has C levels, and so what we're running here is an $R \times C$ factorial ANOVA.

Now that we've got our notation straight, we can compute the sum of squares values for each of the two factors in a relatively familiar way. For Factor A, our between group sum of squares is calculated by assessing the extent to which the (row) marginal means $\bar{Y}_{1..}$, $\bar{Y}_{2..}$ etc, are different from the grand mean $\bar{Y}_{..}$. We do this in the same way that we did for one-way ANOVA: calculate the sum of squared difference between the $\bar{Y}_{i..}$ values and the $\bar{Y}_{..}$ values. Specifically, if there are N people in each group, then we calculate this:

$$SS_A = (N \times C) \sum_{r=1}^R (\bar{Y}_{r..} - \bar{Y}_{..})^2$$

As with one-way ANOVA, the most interesting³ part of this formula is the $(\bar{Y}_{r..} - \bar{Y}_{..})^2$ bit, which corresponds to the squared deviation associated with level r . All that this formula does is calculate this squared deviation for all R levels of the factor, add them up, and then multiply the result by $N \times C$. The reason for this last part is that there are multiple cells in our design that have level r on Factor A: in fact, there are C of them, one corresponding to each possible level of Factor B! For instance, in our toy example, there are *two* different cells in the design corresponding to the `anxitfree` drug: one for people with `no.therapy`, and one for the `CBT` group. Not only that, within each of these cells there are N observations. So, if we want to convert our SS value into a quantity that calculates the between-groups sum of squares on a “per observation” basis, we have to multiply by $N \times C$. The formula for factor B is of course the same thing, just with some subscripts shuffled around:

$$SS_B = (N \times R) \sum_{c=1}^C (\bar{Y}_{..c} - \bar{Y}_{..})^2$$

Now that we have these formulas, we can check them against the R output from the earlier section. First, notice that we calculated all the marginal means (i.e., row marginal means $\bar{Y}_{r..}$ and column marginal means $\bar{Y}_{..c}$) earlier using `aggregate()`, and we also calculated the grand mean. Let's repeat those calculations, but this time we'll save the results to variables so that we can use them in subsequent calculations:

```
> drug.means <- aggregate( mood.gain ~ drug, clin.trial, mean )[,2]
> therapy.means <- aggregate( mood.gain ~ therapy, clin.trial, mean )[,2]
> grand.mean <- mean( clin.trial$mood.gain )
```

³English translation: “least tedious”.

Okay, now let's calculate the sum of squares associated with the main effect of `drug`. There are a total of $N = 3$ people in each group, and $C = 2$ different types of therapy. Or, to put it another way, there are $3 \times 2 = 6$ people who received any particular drug. So our calculations are:

```
> SS.drug <- (3*2) * sum( (drug.means - grand.mean)^2 )
> SS.drug
[1] 3.453333
```

Not surprisingly, this is the same number that you get when you look up the SS value for the drugs factor in the ANOVA table that I presented earlier. We can repeat the same kind of calculation for the effect of therapy. Again there are $N = 3$ people in each group, but since there are $R = 3$ different drugs, this time around we note that there are $3 \times 3 = 9$ people who received CBT, and an additional 9 people who received the placebo. So our calculation is now:

```
> SS.therapy <- (3*3) * sum( (therapy.means - grand.mean)^2 )
> SS.therapy
[1] 0.4672222
```

and we are, once again, unsurprised to see that our calculations are identical to the ANOVA output.

So that's how you calculate the SS values for the two main effects. These SS values are analogous to the between-group sum of squares values that we calculated when doing one-way ANOVA in Chapter 14. However, it's not a good idea to think of them as between-groups SS values anymore, just because we have two different grouping variables and it's easy to get confused. In order to construct an F test, however, we also need to calculate the within-groups sum of squares. In keeping with the terminology that we used in the regression chapter (Chapter 15) and the terminology that R uses when printing out the ANOVA table, I'll start referring to the within-groups SS value as the *residual* sum of squares SS_R .

The easiest way to think about the residual SS values in this context, I think, is to think of it as the leftover variation in the outcome variable after you take into account the differences in the marginal means (i.e., after you remove SS_A and SS_B). What I mean by that is we can start by calculating the total sum of squares, which I'll label SS_T . The formula for this is pretty much the same as it was for one-way ANOVA: we take the difference between each observation Y_{rci} and the grand mean $\bar{Y}_{..}$, square the differences, and add them all up

$$SS_T = \sum_{r=1}^R \sum_{c=1}^C \sum_{i=1}^N (Y_{rci} - \bar{Y}_{..})^2$$

The “triple summation” here looks more complicated than it is. In the first two summations, we're summing across all levels of Factor A (i.e., over all possible rows r in our table), across all levels of Factor B (i.e., all possible columns c). Each rc combination corresponds to a single group, and each group contains N people: so we have to sum across all those people (i.e., all i values) too. In other words, all we're doing here is summing across all observations in the data set (i.e., all possible rci combinations).

At this point, we know the total variability of the outcome variable SS_T , and we know how much of that variability can be attributed to Factor A (SS_A) and how much of it can be attributed to Factor B (SS_B). The residual sum of squares is thus defined to be the variability in Y that *can't* be attributed to either of our two factors. In other words:

$$SS_R = SS_T - (SS_A + SS_B)$$

Of course, there is a formula that you can use to calculate the residual SS directly, but I think that it makes more conceptual sense to think of it like this. The whole point of calling it a residual is that it's the leftover variation, and the formula above makes that clear. I should also note that, in keeping with the terminology used in the regression chapter, it is commonplace to refer to $SS_A + SS_B$ as the variance

attributable to the “ANOVA model”, denoted SS_M , and so we often say that the total sum of squares is equal to the model sum of squares plus the residual sum of squares. Later on in this chapter we’ll see that this isn’t just a surface similarity: ANOVA and regression are actually the same thing under the hood.

In any case, it’s probably worth taking a moment to check that we can calculate SS_R using this formula, and verify that we do obtain the same answer that R produces in its ANOVA table. The calculations are pretty straightforward. First we calculate the total sum of squares:

```
> SS.tot <- sum( (clin.trial$mood.gain - grand.mean)^2 )
> SS.tot
[1] 4.845
```

and then we use it to calculate the residual sum of squares:

```
> SS.res <- SS.tot - (SS.drug + SS.therapy)
> SS.res
[1] 0.9244444
```

Yet again, we get the same answer.

16.1.4 What are our degrees of freedom?

The degrees of freedom are calculated in much the same way as for one-way ANOVA. For any given factor, the degrees of freedom is equal to the number of levels minus 1 (i.e., $R - 1$ for the row variable, Factor A, and $C - 1$ for the column variable, Factor B). So, for the `drugs` factor we obtain $df = 2$, and for the `therapy` factor we obtain $df = 1$. Later on on, when we discuss the interpretation of ANOVA as a regression model (see Section 16.6) I’ll give a clearer statement of how we arrive at this number, but for the moment we can use the simple definition of degrees of freedom, namely that the degrees of freedom equals the number of quantities that are observed, minus the number of constraints. So, for the `drugs` factor, we observe 3 separate group means, but these are constrained by 1 grand mean; and therefore the degrees of freedom is 2. For the residuals, the logic is similar, but not quite the same. The total number of observations in our experiment is 18. The constraints correspond to the 1 grand mean, the 2 additional group means that the `drug` factor introduces, and the 1 additional group mean that the the `therapy` factor introduces, and so our degrees of freedom is 14. As a formula, this is $N - 1 - (R - 1) - (C - 1)$, which simplifies to $N - R - C + 1$.

16.1.5 Factorial ANOVA versus one-way ANOVAs

Now that we’ve seen *how* a factorial ANOVA works, it’s worth taking a moment to compare it to the results of the one way analyses, because this will give us a really good sense of *why* it’s a good idea to run the factorial ANOVA. In Chapter 14 that I ran a one-way ANOVA that looked to see if there are any differences between drugs, and a second one-way ANOVA to see if there were any differences between therapies. As we saw in Section 16.1.1, the null and alternative hypotheses tested by the one-way ANOVAs are in fact identical to the hypotheses tested by the factorial ANOVA. Looking even more carefully at the ANOVA tables, we can see that the sum of squares associated with the factors are identical in the two different analyses (3.45 for `drug` and 0.92 for `therapy`), as are the degrees of freedom (2 for `drug`, 1 for `therapy`). But they don’t give the same answers! Most notably, when we ran the one-way ANOVA for `therapy` in Section 14.11 we didn’t find a significant effect (the p -value was 0.21). However, when we look at the main effect of `therapy` within the context of the two-way ANOVA, we do get a significant effect ($p = .019$). The two analyses are clearly not the same.

Why does that happen? The answer lies in understanding how the *residuals* are calculated. Recall that the whole idea behind an *F*-test is to compare the variability that can be attributed to a particular factor with the variability that cannot be accounted for (the residuals). If you run a one-way ANOVA for `therapy`, and therefore ignore the effect of `drug`, the ANOVA will end up dumping all of the drug-induced variability into the residuals! This has the effect of making the data look more noisy than they really are, and the effect of `therapy` which is correctly found to be significant in the two-way ANOVA now becomes non-significant. If we ignore something that actually matters (e.g., `drug`) when trying to assess the contribution of something else (e.g., `therapy`) then our analysis will be distorted. Of course, it's perfectly okay to ignore variables that are genuinely irrelevant to the phenomenon of interest: if we had recorded the colour of the walls, and that turned out to be non-significant in a three-way ANOVA (i.e. `mood.gain ~ drug + therapy + wall.colour`), it would be perfectly okay to disregard it and just report the simpler two-way ANOVA that doesn't include this irrelevant factor. What you shouldn't do is drop variables that actually make a difference!

16.1.6 What kinds of outcomes does this analysis capture?

The ANOVA model that we've been talking about so far covers a range of different patterns that we might observe in our data. For instance, in a two-way ANOVA design, there are four possibilities: (a) only Factor A matters, (b) only Factor B matters, (c) both A and B matter, and (d) neither A nor B matters. An example of each of these four possibilities is plotted in Figure 16.1.

16.2

Factorial ANOVA 2: balanced designs, interactions allowed

The four patterns of data shown in Figure 16.1 are all quite realistic: there are a great many data sets that produce exactly those patterns. However, they are not the whole story, and the ANOVA model that we have been talking about up to this point is not sufficient to fully account for a table of group means. Why not? Well, so far we have the ability to talk about the idea that drugs can influence mood, and therapy can influence mood, but no way of talking about the possibility of an `interaction` between the two. An interaction between A and B is said to occur whenever the effect of Factor A is *different*, depending on which level of Factor B we're talking about. Several examples of an interaction effect with the context of a 2×2 ANOVA are shown in Figure 16.2. To give a more concrete example, suppose that the operation of Anxitriptal and Joyzepam is governed quite different physiological mechanisms, and one consequence of this is that while Joyzepam has more or less the same effect on mood regardless of whether one is in therapy, Anxitriptal is actually much more effective when administered in conjunction with CBT. The ANOVA that we developed in the previous section does not capture this idea. To get some idea of whether an interaction is actually happening here, it helps to plot the various group means. There are quite a few different ways draw these plots in R. One easy way is to use the `interaction.plot()` function, but this function won't draw error bars for you. A fairly simple function that will include error bars for you is the `lineplot.CI()` function in the `sciplots` package (see Section 10.5.2). The command

```
> lineplot.CI( x.factor = clin.trial$drug,
+               response = clin.trial$mood.gain,
+               group = clin.trial$therapy,
+               ci.fun = ciMean,
+               xlab = "drug",
+               ylab = "mood gain" )
```

produces the output is shown in Figure 16.3 (don't forget that the `ciMean` function is in the `lsr` package, so you need to have `lsr` loaded!). Our main concern relates to the fact that the two lines aren't parallel.

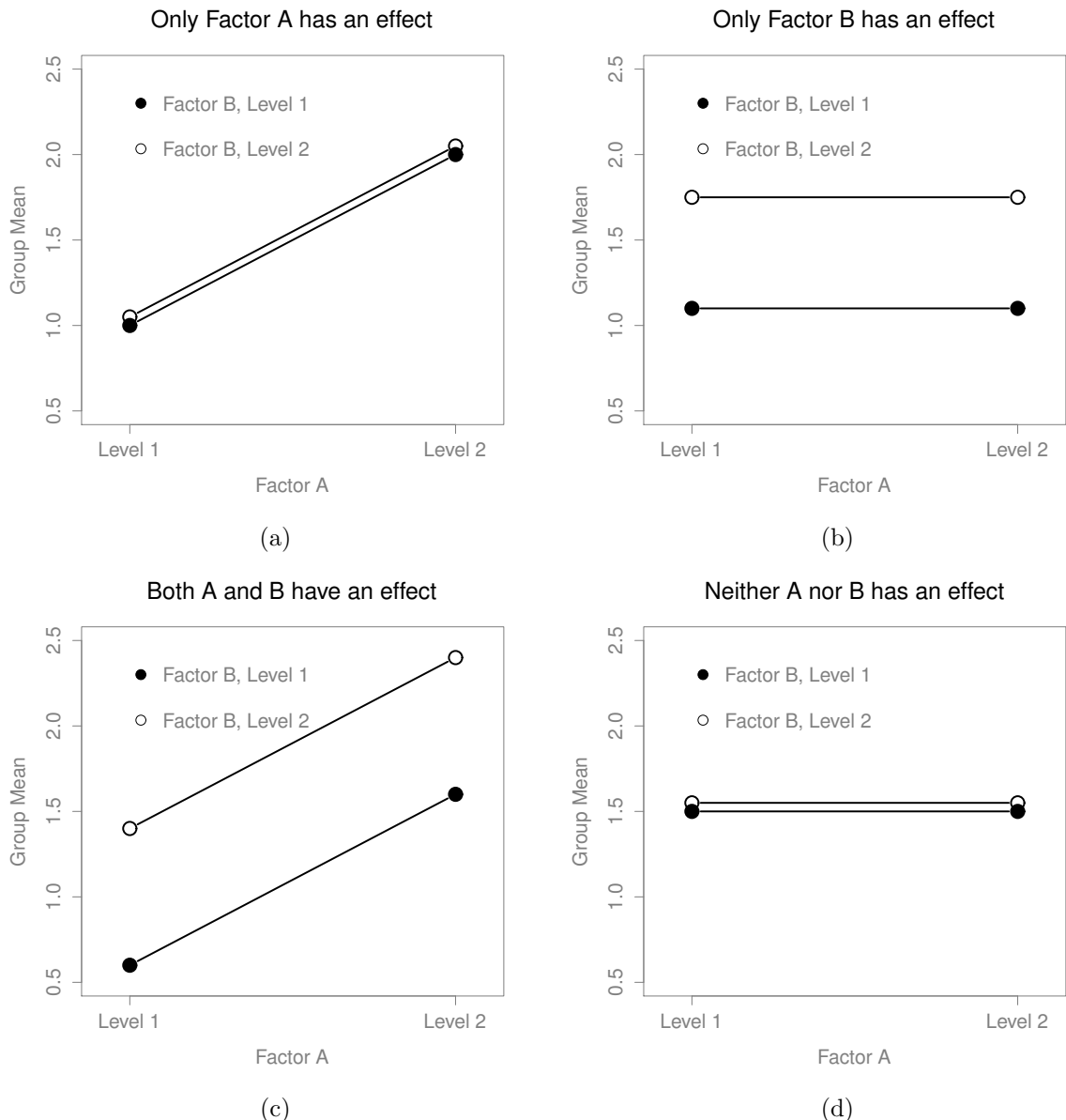


Figure 16.1: The four different outcomes for a 2×2 ANOVA when no interactions are present. In panel (a) we see a main effect of Factor A, and no effect of Factor B. Panel (b) shows a main effect of Factor B but no effect of Factor A. Panel (c) shows main effects of both Factor A and Factor B. Finally, panel (d) shows no effect of either factor.

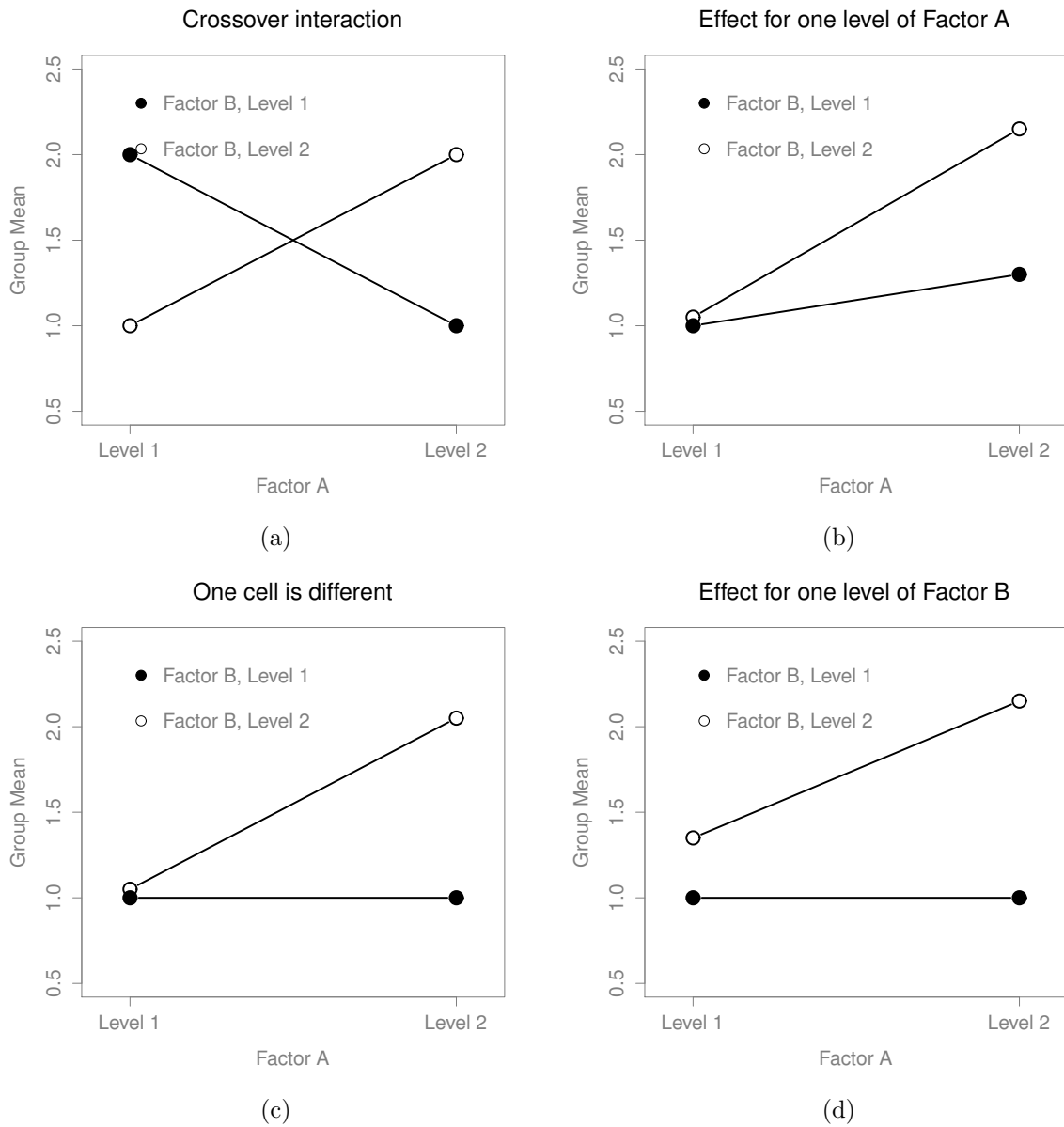


Figure 16.2: Qualitatively different interactions for a 2×2 ANOVA

The effect of CBT (difference between solid line and dotted line) when the drug is Joyzepam (right side) appears to be near zero, even smaller than the effect of CBT when a placebo is used (left side). However, when Anxitfree is administered, the effect of CBT is larger than the placebo (middle). Is this effect real, or is this just random variation due to chance? Our original ANOVA cannot answer this question, because we make no allowances for the idea that interactions even exist! In this section, we'll fix this problem.

16.2.1 **What exactly is an interaction effect?**

The key idea that we're going to introduce in this section is that of an interaction effect. What that means for our R formulas is that we'll write down models like

```
mood.gain ~ drug + therapy + drug:therapy
```

so although there are only two *factors* involved in our model (i.e., `drug` and `therapy`), there are three distinct **terms** (i.e., `drug`, `therapy` and `drug:therapy`). That is, in addition to the main effects of `drug` and `therapy`, we have a new component to the model, which is our interaction term `drug:therapy`. Intuitively, the idea behind an interaction effect is fairly simple: it just means that the effect of Factor A is different, depending on which level of Factor B we're talking about. But what does that actually mean in terms of our data? Figure 16.2 depicts several different patterns that, although quite different to each other, would all count as an interaction effect. So it's not entirely straightforward to translate this qualitative idea into something mathematical that a statistician can work with. As a consequence, the way that the idea of an interaction effect is formalised in terms of null and alternative hypotheses is slightly difficult, and I'm guessing that a lot of readers of this book probably won't be all that interested. Even so, I'll try to give the basic idea here.

To start with, we need to be a little more explicit about our main effects. Consider the main effect of Factor A (`drug` in our running example). We originally formulated this in terms of the null hypothesis that the two marginal means $\mu_{r\cdot}$ are all equal to each other. Obviously, if all of these are equal to each other, then they must also be equal to the grand mean $\mu_{..}$ as well, right? So what we can do is define the *effect* of Factor A at level r to be equal to the difference between the marginal mean $\mu_{r\cdot}$ and the grand mean $\mu_{..}$. Let's denote this effect by α_r , and note that

$$\alpha_r = \mu_{r\cdot} - \mu_{..}$$

Now, by definition all of the α_r values must sum to zero, for the same reason that the average of the marginal means $\mu_{r\cdot}$ must be the grand mean $\mu_{..}$. We can similarly define the effect of Factor B at level i to be the difference between the column marginal mean $\mu_{.c}$ and the grand mean $\mu_{..}$.

$$\beta_c = \mu_{.c} - \mu_{..}$$

and once again, these β_c values must sum to zero. The reason that statisticians sometimes like to talk about the main effects in terms of these α_r and β_c values is that it allows them to be precise about what it means to say that there is no interaction effect. If there is no interaction at all, then these α_r and β_c values will perfectly describe the group means μ_{rc} . Specifically, it means that

$$\mu_{rc} = \mu_{..} + \alpha_r + \beta_c$$

That is, there's nothing *special* about the group means that you couldn't predict perfectly by knowing all the marginal means. And that's our null hypothesis, right there. The alternative hypothesis is that

$$\mu_{ij} \neq \mu_{..} + \alpha_r + \beta_c$$

for at least one group rc in our table. However, statisticians often like to write this slightly differently. They'll usually define the specific interaction associated with group rc to be some number, awkwardly

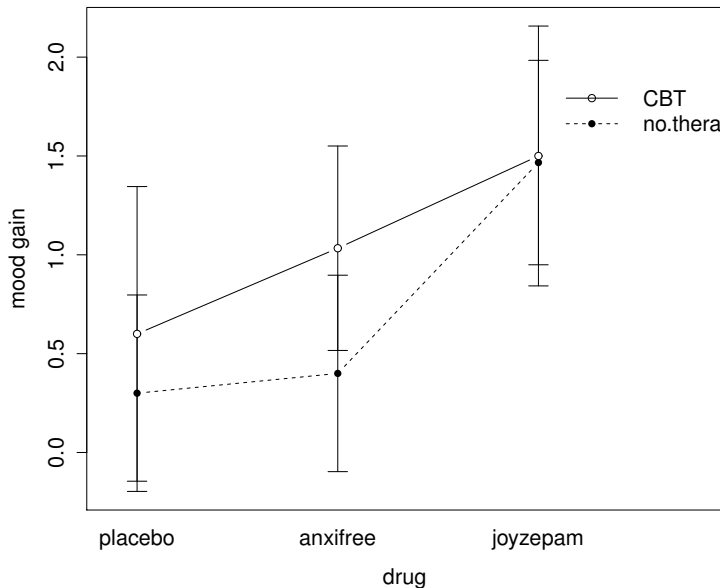


Figure 16.3: An interaction plot for the group means in the clinical trial data. The command to produce it is included in the main text. You'll notice that the legend doesn't quite fit properly. You can fix this by playing around with the `x.leg` and `y.leg` arguments: type `?lineplot.CI` for details.

.....

referred to as $(\alpha\beta)_{rc}$, and then they will say that the alternative hypothesis is that

$$\mu_{rc} = \mu_{..} + \alpha_r + \beta_c + (\alpha\beta)_{rc}$$

where $(\alpha\beta)_{rc}$ is non-zero for at least one group. This notation is kind of ugly to look at, but it is handy as we'll see in the next section when discussing how to calculate the sum of squares.

16.2.2 Calculating sums of squares for the interaction

How should we calculate the sum of squares for the interaction terms, $SS_{A:B}$? Well, first off, it helps to notice how the previous section defined the interaction effect in terms of the extent to which the actual group means differ from what you'd expect by just looking at the marginal means. Of course, all of those formulas refer to population parameters rather than sample statistics, so we don't actually know what they are. However, we can estimate them by using sample means in place of population means. So for Factor A, a good way to estimate the main effect at level r as the difference between the *sample* marginal mean \bar{Y}_{rc} and the sample grand mean $\bar{Y}_{..}$. That is, we would use this as our estimate of the effect:

$$\hat{\alpha}_r = \bar{Y}_{r.} - \bar{Y}_{..}$$

Similarly, our estimate of the main effect of Factor B at level c can be defined as follows:

$$\hat{\beta}_c = \bar{Y}_{.c} - \bar{Y}_{..}$$

Now, if you go back to the formulas that I used to describe the SS values for the two main effects, you'll notice that these effect terms are exactly the quantities that we were squaring and summing! So what's the analog of this for interaction terms? The answer to this can be found by first rearranging the formula for the group means μ_{rc} under the alternative hypothesis, so that we get this:

$$\begin{aligned} (\alpha\beta)_{rc} &= \mu_{rc} - \mu_{..} - \alpha_r - \beta_c \\ &= \mu_{rc} - \mu_{..} - (\mu_{r.} - \mu_{..}) - (\mu_{.c} - \mu_{..}) \\ &= \mu_{rc} - \mu_{r.} - \mu_{.c} + \mu_{..} \end{aligned}$$

So, once again, if we substitute our sample statistics in place of the population means, we get the following as our estimate of the interaction effect for group rc , which is

$$(\hat{\alpha}\beta)_{rc} = \bar{Y}_{rc} - \bar{Y}_{r.} - \bar{Y}_{.c} + \bar{Y}_{..}$$

Now all we have to do is sum all of these estimates across all R levels of Factor A and all C levels of Factor B, and we obtain the following formula for the sum of squares associated with the interaction as a whole:

$$SS_{A:B} = N \sum_{r=1}^R \sum_{c=1}^C (\bar{Y}_{rc} - \bar{Y}_{r.} - \bar{Y}_{.c} + \bar{Y}_{..})^2$$

where, we multiply by N because there are N observations in each of the groups, and we want our SS values to reflect the variation among *observations* accounted for by the interaction, not the variation among groups.

Now that we have a formula for calculating $SS_{A:B}$, it's important to recognise that the interaction term is part of the model (of course), so the total sum of squares associated with the model, SS_M is now equal to the sum of the three relevant SS values, $SS_A + SS_B + SS_{A:B}$. The residual sum of squares SS_R is still defined as the leftover variation, namely $SS_T - SS_M$, but now that we have the interaction term this becomes

$$SS_R = SS_T - (SS_A + SS_B + SS_{A:B})$$

As a consequence, the residual sum of squares SS_R will be smaller than in our original ANOVA that didn't include interactions.

16.2.3 Degrees of freedom for the interaction

Calculating the degrees of freedom for the interaction is, once again, slightly trickier than the corresponding calculation for the main effects. To start with, let's think about the ANOVA model as a whole. Once we include interaction effects in the model, we're allowing every single group has a unique mean, μ_{rc} . For an $R \times C$ factorial ANOVA, this means that there are $R \times C$ quantities of interest in the model, and only the one constraint: all of the group means need to average out to the grand mean. So the model as a whole needs to have $(R \times C) - 1$ degrees of freedom. But the main effect of Factor A has $R - 1$ degrees of freedom, and the main effect of Factor B has $C - 1$ degrees of freedom. Which means that the degrees of freedom associated with the interaction is

$$\begin{aligned} df_{A:B} &= (R \times C - 1) - (R - 1) - (C - 1) \\ &= RC - R - C + 1 \\ &= (R - 1)(C - 1) \end{aligned}$$

which is just the product of the degrees of freedom associated with the row factor and the column factor.

What about the residual degrees of freedom? Because we've added interaction terms, which absorb some degrees of freedom, there are fewer residual degrees of freedom left over. Specifically, note that if

the model with interaction has a total of $(R \times C) - 1$, and there are N observations in your data set that are constrained to satisfy 1 grand mean, your residual degrees of freedom now become $N - (R \times C) - 1 + 1$, or just $N - (R \times C)$.

16.2.4 Running the ANOVA in R

Adding interaction terms to the ANOVA model in R is straightforward. Returning to our running example of the clinical trial, in addition to the main effect terms of `drug` and `therapy`, we include the interaction term `drug:therapy`. So the R command to create the ANOVA model now looks like this:

```
> model.3 <- aov( mood.gain ~ drug + therapy + drug:therapy, clin.trial )
```

However, R allows a convenient shorthand. Instead of typing out all three terms, you can shorten the right hand side of the formula to `drug*therapy`. The `*` operator inside the formula is taken to indicate that you want both main effects and the interaction. So we can also run our ANOVA like this, and get the same answer:

```
> model.3 <- aov( mood.gain ~ drug * therapy, clin.trial )
> summary( model.3 )
   Df Sum Sq Mean Sq F value    Pr(>F)
drug        2   3.45   1.727   31.71 1.6e-05 ***
therapy      1   0.47   0.467    8.58   0.013 *
drug:therapy 2   0.27   0.136    2.49   0.125
Residuals   12   0.65   0.054
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

As it turns out, while we do have a significant main effect of drug ($F_{2,12} = 31.7, p < .001$) and therapy type ($F_{1,12} = 8.6, p = .013$), there is no significant interaction between the two ($F_{2,12} = 2.5, p = 0.125$).

16.2.5 Interpreting the results

There's a couple of very important things to consider when interpreting the results of factorial ANOVA. Firstly, there's the same issue that we had with one-way ANOVA, which is that if you obtain a significant main effect of (say) `drug`, it doesn't tell you anything about which drugs are different to one another. To find that out, you need to run additional analyses. We'll talk about some analyses that you can run in Sections 16.7 and 16.8. The same is true for interaction effects: knowing that there's a significant interaction doesn't tell you anything about what kind of interaction exists. Again, you'll need to run additional analyses.

Secondly, there's a very peculiar interpretation issue that arises when you obtain a significant interaction effect but no corresponding main effect. This happens sometimes. For instance, in the crossover interaction shown in Figure 16.2a, this is exactly what you'd find: in this case, neither of the main effects would be significant, but the interaction effect would be. This is a difficult situation to interpret, and people often get a bit confused about it. The general advice that statisticians like to give in this situation is that you shouldn't pay much attention to the main effects when an interaction is present. The reason they say this is that, although the tests of the main effects are perfectly valid from a mathematical point of view, when there is a significant interaction effect the main effects rarely test interesting hypotheses. Recall from Section 16.1.1 that the null hypothesis for a main effect is that the *marginal means* are equal to each other, and that a marginal mean is formed by averaging across several different groups. But if you have a significant interaction effect, then you *know* that the groups that comprise the marginal mean aren't homogeneous, so it's not really obvious why you would care about those marginal means.

Here's what I mean. Again, let's stick with a clinical example. Suppose that we had a 2×2 design comparing two different treatments for phobias (e.g., systematic desensitisation vs flooding), and two different anxiety reducing drugs (e.g., Anxitriptaline vs Sertraline). Now suppose what we found was that Anxitriptaline had no effect when desensitisation was the treatment, and Sertraline had no effect when flooding was the treatment. But both were pretty effective for the other treatment. This is a classic crossover interaction, and what we'd find when running the ANOVA is that there is no main effect of drug, but a significant interaction. Now, what does it actually *mean* to say that there's no main effect? Well, it means that, if we average over the two different psychological treatments, then the *average* effect of Anxitriptaline and Sertraline is the same. But why would anyone care about that? When treating someone for phobias, it is never the case that a person can be treated using an "average" of flooding and desensitisation: that doesn't make a lot of sense. You either get one or the other. For one treatment, one drug is effective; and for the other treatment, the other drug is effective. The interaction is the important thing; the main effect is kind of irrelevant.

This sort of thing happens a lot: the main effect are tests of marginal means, and when an interaction is present we often find ourselves not being terribly interested in marginal means, because they imply averaging over things that the interaction tells us shouldn't be averaged! Of course, it's not always the case that a main effect is meaningless when an interaction is present. Often you can get a big main effect and a very small interaction, in which case you can still say things like "drug A is generally more effective than drug B" (because there was a big effect of drug), but you'd need to modify it a bit by adding that "the difference in effectiveness was different for different psychological treatments". In any case, the main point here is that whenever you get a significant interaction you should stop and *think* about what the main effect actually means in this context. Don't automatically assume that the main effect is interesting.

16.3

Effect size, estimated means, and confidence intervals

In this section I'll discuss a few additional quantities that you might find yourself wanting to calculate for a factorial ANOVA. The main thing you will probably want to calculate is the effect size for each term in your model, but you may also want to R to give you some estimates for the group means and associated confidence intervals.

16.3.1 **Effect sizes**

The effect size calculations for a factorial ANOVA is pretty similar to those used in one way ANOVA (see Section 14.4). Specifically, we can use η^2 (eta-squared) as simple way to measure how big the overall effect is for any particular term. As before, η^2 is defined by dividing the sum of squares associated with that term by the total sum of squares. For instance, to determine the size of the main effect of Factor A, we would use the following formula

$$\eta_A^2 = \frac{SS_A}{SS_T}$$

As before, this can be interpreted in much the same way as R^2 in regression.⁴ It tells you the proportion of variance in the outcome variable that can be accounted for by the main effect of Factor A. It is therefore a number that ranges from 0 (no effect at all) to 1 (accounts for *all* of the variability in the

⁴This chapter seems to be setting a new record for the number of different things that the letter R can stand for: so far we have R referring to the software package, the number of rows in our table of means, the residuals in the model, and now the correlation coefficient in a regression. Sorry: we clearly don't have enough letters in the alphabet. However, I've tried pretty hard to be clear on which thing R is referring to in each case.

outcome). Moreover, the sum of all the η^2 values, taken across all the terms in the model, will sum to the total R^2 for the ANOVA model. If, for instance, the ANOVA model fits perfectly (i.e., there is no within-groups variability at all!), the η^2 values will sum to 1. Of course, that rarely if ever happens in real life.

However, when doing a factorial ANOVA, there is a second measure of effect size that people like to report, known as partial η^2 . The idea behind partial η^2 (which is sometimes denoted $p\eta^2$ or η_p^2) is that, when measuring the effect size for a particular term (say, the main effect of Factor A), you want to deliberately ignore the other effects in the model (e.g., the main effect of Factor B). That is, you would pretend that the effect of all these other terms is zero, and then calculate what the η^2 value would have been. This is actually pretty easy to calculate. All you have to do is remove the sum of squares associated with the other terms from the denominator. In other words, if you want the partial η^2 for the main effect of Factor A, the denominator is just the sum of the SS values for Factor A and the residuals:

$$\text{partial } \eta_A^2 = \frac{\text{SS}_A}{\text{SS}_A + \text{SS}_R}$$

This will always give you a larger number than η^2 , which the cynic in me suspects accounts for the popularity of partial η^2 . And once again you get a number between 0 and 1, where 0 represents no effect. However, it's slightly trickier to interpret what a large partial η^2 value means. In particular, you can't actually compare the partial η^2 values across terms! Suppose, for instance, there is no within-groups variability at all: if so, $\text{SS}_R = 0$. What that means is that *every* term has a partial η^2 value of 1. But that doesn't mean that all terms in your model are equally important, or indeed that they are equally large. All it means is that all terms in your model have effect sizes that are large *relative to the residual variation*. It is not comparable across terms.

To see what I mean by this, it's useful to see a concrete example. Once again, we'll use the `etaSquared()` function from the `lsr` package. As before, we input the `aov` object for which we want the η^2 calculations performed, and R outputs a matrix showing the effect sizes for each term in the model. First, let's have a look at the effect sizes for the original ANOVA without the interaction term:

```
> etaSquared( model.2 )
      eta.sq partial.eta.sq
drug      0.712762     0.78883
therapy   0.096434     0.33573
```

Looking at the η^2 values first, we see that `drug` accounts for 71.3% of the variance (i.e. $\eta^2 = 0.713$) in `mood.gain`, whereas `therapy` only accounts for 9.6%. This leaves a total of 19.1% of the variation unaccounted for (i.e., the residuals constitute 19.1% of the variation in the outcome). Overall, this implies that we have a very large effect⁵ of `drug` and a modest effect of `therapy`.

Now let's look at the partial η^2 values. Because the effect of `therapy` isn't all that large, controlling for it doesn't make much of a difference, so the partial η^2 for `drug` doesn't increase very much, and we obtain a value of $p\eta^2 = 0.789$. In contrast, because the effect of `drug` was very large, controlling for it makes a big difference, and so when we calculate the partial η^2 for `therapy` you can see that it rises to $p\eta^2 = 0.336$. The question that we have to ask ourselves is, what does these partial η^2 values actually *mean*? The way I generally interpret the partial η^2 for the main effect of Factor A is to interpret it as a statement about a hypothetical experiment in which *only* Factor A was being varied. So, even though in *this* experiment we varied both A and B, we can easily imagine an experiment in which only Factor A was varied: the partial η^2 statistic tells you how much of the variance in the outcome variable you would expect to see accounted for in that experiment. However, it should be noted that this interpretation – like many things associated with main effects – doesn't make a lot of sense when there is a large and significant interaction effect.

⁵Implausibly large, I would think: the artificiality of this data set is really starting to show!

Speaking of interaction effects, here's what we get when we calculate the effect sizes for the model that includes the interaction term. As you can see, the η^2 values for the main effects don't change, but the partial η^2 values do:

```
> etaSquared( model.3 )
      eta.sq partial.eta.sq
drug      0.712762    0.84091
therapy   0.096434    0.41696
drug:therapy 0.055957  0.29327
```

16.3.2 Estimated group means

In many situations you will find yourself wanting to report estimates of all the group means based on the results of your ANOVA, as well as confidence intervals associated with them. You can use the `effect()` function in the `effects` package to do this (don't forget to install the package if you don't have it already!). If the ANOVA that you have run is a **saturated model** (i.e., contains all possible main effects and all possible interaction effects) then the estimates of the group means are actually identical to the sample means, though the confidence intervals will use a pooled estimate of the standard errors, rather than use a separate one for each group. To illustrate this, let's apply the `effect()` function to our saturated model (i.e., `model.3`) for the clinical trial data. The `effect()` function contains two arguments we care about: the `term` argument specifies what terms in the model we want the means to be calculated for, and the `mod` argument specifies the model:

```
> library(effects)
> eff <- effect( term = "drug*therapy", mod = model.3 )
> eff

drug*therapy effect
  therapy
drug      no.therapy      CBT
placebo   0.300000 0.600000
anxifree  0.400000 1.033333
joyzepam   1.466667 1.500000
```

Notice that these are actually the same numbers we got when computing the sample means earlier (i.e., the `group.means` variable that we computed using `aggregate()`). One useful thing that we can do using the effect variable `eff`, however, is extract the confidence intervals using the `summary()` function:

```
> summary(eff)

drug*therapy effect
  therapy
drug      no.therapy      CBT
placebo   0.300000 0.600000
anxifree  0.400000 1.033333
joyzepam   1.466667 1.500000

Lower 95 Percent Confidence Limits
  therapy
drug      no.therapy      CBT
placebo  0.006481093 0.3064811
anxifree 0.106481093 0.7398144
joyzepam 1.173147759 1.2064811
```

```

Upper 95 Percent Confidence Limits
      therapy
drug      no.therapy      CBT
placebo   0.5935189 0.8935189
anxifree  0.6935189 1.3268522
joyzepam  1.7601856 1.7935189

```

In this output, we see that the estimated mean mood gain for the placebo group with no therapy was 0.300, with a 95% confidence interval from 0.006 to 0.594. Note that these are not the same confidence intervals that you would get if you calculated them separately for each group, because of the fact that the ANOVA model assumes homogeneity of variance and therefore uses a pooled estimate of the standard deviation.

When the model doesn't contain the interaction term, then the estimated group means will be different from the sample means. Instead of reporting the sample mean, the `effect()` function will calculate the value of the group means that would be expected on the basis of the marginal means (i.e., assuming no interaction). Using the notation we developed earlier, the estimate reported for μ_{rc} , the mean for level r on the (row) Factor A and level c on the (column) Factor B would be $\mu_{..} + \alpha_r + \beta_c$. If there are genuinely no interactions between the two factors, this is actually a better estimate of the population mean than the raw sample mean would be. The command to obtain these estimates is actually identical to the last one, except that we use `model.2`. When you do this, R will give you a warning message:

```

> eff <- effect( "drug*therapy", model.2 )
Warning message:
In analyze.model(term, mod, xlevels, default.levels) :
  drug:therapy does not appear in the model

```

but this isn't anything to worry about. This is R being polite, and letting you know that the estimates it is constructing are based on the assumption that no interactions exist. It kind of makes sense that it would do this: when we use "`drug*therapy`" as our input, we're telling R that we want it to output the estimated group means (rather than marginal means), but the actual input "`drug*therapy`" might mean that you want interactions included or you might not. There's no *actual* ambiguity here, because the model itself either does or doesn't have interactions, but the authors of the function thought it sensible to include a warning just to make sure that you've specified the actual model you care about. But, assuming that we genuinely don't believe that there are any interactions, `model.2` is the right model to use, so we can ignore this warning.⁶ In any case, when we inspect the output, we get the following table of estimated group means:

```

> eff

drug*therapy effect
      therapy
drug      no.therapy      CBT
placebo   0.2888889 0.6111111
anxifree  0.5555556 0.8777778
joyzepam  1.3222222 1.6444444

```

As before, we can obtain confidence intervals using the following command:

```
> summary( eff )
```

but the output looks pretty much the same as last time, and this book is already way too long, so I won't include it here.

⁶In fact, there's a function `Effect()` within the `effects` package that has slightly different arguments, but computes the same things, and won't give you this warning message.

16.4

Assumption checking

As with one-way ANOVA, the key assumptions of factorial ANOVA are homogeneity of variance (all groups have the same standard deviation), normality of the residuals, and independence of the observations. The first two are things we can test for. The third is something that you need to assess yourself by asking if there are any special relationships between different observations. Additionally, if you aren't using a saturated model (e.g., if you've omitted the interaction terms) then you're also assuming that the omitted terms aren't important. Of course, you can check this last one by running an ANOVA with the omitted terms included and see if they're significant, so that's pretty easy. What about homogeneity of variance and normality of the residuals? As it turns out, these are pretty easy to check: it's no different to the checks we did for a one-way ANOVA.

16.4.1 Levene test for homogeneity of variance

To test whether the groups have the same variance, we can use the Levene test. The theory behind the Levene test was discussed in Section 14.7, so I won't discuss it again. Once again, you can use the `leveneTest()` function in the `car` package to do this. This function expects that you have a saturated model (i.e., included all of the relevant terms), because the test is primarily concerned with the within-group variance, and it doesn't really make a lot of sense to calculate this any way other than with respect to the full model. So we try either of the following commands:

```
> leveneTest( model.2 )
> leveneTest( mood.gain ~ drug + therapy, clin.trial )
```

R will spit out the following error:

```
Error in leveneTest.formula(formula(y), data = model.frame(y), ...):
  Model must be completely crossed formula only.
```

Instead, if you want to run the Levene test, you need to specify a saturated model. Either of the following two commands would work:⁷

```
> leveneTest( model.3 )
> leveneTest( mood.gain ~ drug * therapy, clin.trial )
```

and would produce the following as the output:

```
Levene's Test for Homogeneity of Variance (center = median)
  Df F value Pr(>F)
group  5  0.0955 0.9912
      12
```

The fact that the Levene test is non-significant means that we can safely assume that the homogeneity of variance assumption is not violated.

⁷Due to the way that the `leveneTest()` function is implemented, however, if you use a formula like `mood.gain ~ drug + therapy + drug:therapy`, or input an ANOVA object based on a formula like this, you actually get the error message. That shouldn't happen, because this actually is a fully crossed model. However, there's a quirky shortcut in the way that the `leveneTest()` function checks whether your model is fully crossed that means that it doesn't recognise this as a fully crossed model. Essentially what the function is doing is checking that you used `*` (which ensures that the model is fully crossed), and not `+` or `:` in your model formula. So if you've manually typed out all of the relevant terms for a fully crossed model, the `leveneTest()` function doesn't detect it. I think this is a bug.

16.4.2 Normality of residuals

As with one-way ANOVA, we can test for the normality of residuals in a straightforward fashion (see Section 14.9). First, we use the `residuals()` function to extract the residuals from the model itself, and then we can examine those residuals in a few different ways. It's generally a good idea to examine them graphically, by drawing histograms (i.e., `hist()` function) and QQ plots (i.e., `qqnorm()` function). If you want a formal test for the normality of the residuals, then we can run the Shapiro-Wilk test (i.e., `shapiro.test()`). If we wanted to check the residuals with respect to `model.2` (i.e., the model with both main effects but no interactions) then we could do the following:

```
> resid <- residuals( model.2 ) # pull the residuals
> hist( resid ) # draw a histogram
> qqnorm( resid ) # draw a normal QQ plot
> shapiro.test( resid ) # run the Shapiro-Wilk test

Shapiro-Wilk normality test

data: resid
W = 0.9563, p-value = 0.5329
```

I haven't included the plots (you can draw them yourself if you want to see them), but you can see from the non-significance of the Shapiro-Wilk test that normality isn't violated here.

16.5

The *F* test as a model comparison

At this point, I want to talk in a little more detail about what the *F*-tests in an ANOVA are actually doing. In the context of ANOVA, I've been referring to the *F*-test as a way of testing whether a particular term in the model (e.g., main effect of Factor A) is significant. This interpretation is perfectly valid, but it's not necessarily the most useful way to think about the test. In fact, it's actually a fairly limiting way of thinking about what the *F*-test does. Consider the clinical trial data we've been working with in this chapter. Suppose I want to see if there are *any* effects of any kind that involve `therapy`. I'm not fussy: I don't care if it's a main effect or an interaction effect.⁸ One thing I could do is look at the output for `model.3` earlier: in this model we did see a main effect of therapy ($p = .013$) but we did not see an interaction effect ($p = .125$). That's kind of telling us what we want to know, but it's not quite the same thing. What we really want is a single test that *jointly* checks the main effect of therapy and the interaction effect.

Given the way that I've been describing the ANOVA *F*-test up to this point, you'd be tempted to think that this isn't possible. On the other hand, if you recall the chapter on regression (in Section 15.9), we were able to use *F*-tests to make comparisons between a wide variety of regression models. Perhaps something of that sort is possible with ANOVA? And of course, the answer here is yes. The thing that you really need to understand is that the *F*-test, as it is used in both ANOVA and regression, is really a comparison of *two* statistical models. One of these models is the full model (alternative hypothesis), and the other model is a simpler model that is missing one or more of the terms that the full model includes (null hypothesis). The null model cannot contain any terms that are not in the full model. In the example I gave above, the full model is `model.3`, and it contains a main effect for therapy, a main effect for drug, and the drug by therapy interaction term. The null model would be `model.1` since it contains only the main effect of drug.

⁸There could be all sorts of reasons for doing this, I would imagine.

16.5.1 The F test comparing two models

Let's frame this in a slightly more abstract way. We'll say that our full model can be written as an R formula that contains several different terms, say $\mathbf{Y} \sim \mathbf{A} + \mathbf{B} + \mathbf{C} + \mathbf{D}$. Our null model only contains some subset of these terms, say $\mathbf{Y} \sim \mathbf{A} + \mathbf{B}$. Some of these terms might be main effect terms, others might be interaction terms. It really doesn't matter. The only thing that matters here is that we want to treat some of these terms as the "starting point" (i.e. the terms in the null model, \mathbf{A} and \mathbf{B}), and we want to see if including the other terms (i.e., \mathbf{C} and \mathbf{D}) leads to a significant improvement in model performance, over and above what could be achieved by a model that includes only \mathbf{A} and \mathbf{B} . In essence, we have null and alternative hypotheses that look like this:

Hypothesis	Correct model?	R formula for correct model
Null	M_0	$\mathbf{Y} \sim \mathbf{A} + \mathbf{B}$
Alternative	M_1	$\mathbf{Y} \sim \mathbf{A} + \mathbf{B} + \mathbf{C} + \mathbf{D}$

Is there a way of making this comparison directly?

To answer this, let's go back to fundamentals. As we saw in Chapter 14, the F -test is constructed from two kinds of quantity: sums of squares (SS) and degrees of freedom (df). These two things define a mean square value ($MS = SS/df$), and we obtain our F statistic by contrasting the MS value associated with "the thing we're interested in" (the model) with the MS value associated with "everything else" (the residuals). What we want to do is figure out how to talk about the SS value that is associated with the *difference* between two models. It's actually not all that hard to do.

Let's start with the fundamental rule that we used throughout the chapter on regression:

$$SS_T = SS_M + SS_R$$

That is, the total sums of squares (i.e., the overall variability of the outcome variable) can be decomposed into two parts: the variability associated with the model SS_M , and the residual or leftover variability, SS_R . However, it's kind of useful to rearrange this equation slightly, and say that the SS value associated with a model is defined like this...

$$SS_M = SS_T - SS_R$$

Now, in our scenario, we have two models: the null model (M_0) and the full model (M_1):

$$\begin{aligned} SS_{M_0} &= SS_T - SS_{R_0} \\ SS_{M_1} &= SS_T - SS_{R_1} \end{aligned}$$

Next, let's think about what it is we *actually* care about here. What we're interested in is the *difference* between the full model and the null model. So, if we want to preserve the idea that what we're doing is an "analysis of the variance" (ANOVA) in the outcome variable, what we should do is define the SS associated with the difference to be equal to the difference in the SS :

$$\begin{aligned} SS_\Delta &= SS_{M_1} - SS_{M_0} \\ &= (SS_T - SS_{R_1}) - (SS_T - SS_{R_0}) \\ &= SS_{R_0} - SS_{R_1} \end{aligned}$$

Now that we have our degrees of freedom, we can calculate mean squares and F values in the usual way. Specifically, we're interested in the mean square for the difference between models, and the mean square for the residuals associated with the *full* model (M_1), which are given by

$$MS_\Delta = \frac{SS_\Delta}{df_\Delta}$$

$$MS_{R1} = \frac{SS_{R1}}{df_{R1}}$$

Finally, taking the ratio of these two gives us our F statistic:

$$F = \frac{MS_{\Delta}}{MS_{R1}}$$

16.5.2 Running the test in R

At this point, it may help to go back to our concrete example. The null model here is `model.1`, which stipulates that there is a main effect of drug, but no other effects exist. We expressed this via the model formula `mood.gain ~ drug`. The alternative model here is `model.3`, which stipulates that there is a main effect of drug, a main effect of therapy, *and* an interaction. If we express this in the “long” format, this model corresponds to the formula `mood.gain ~ drug + therapy + drug:therapy`, though we often express this using the `*` shorthand. The key thing here is that if we compare `model.1` to `model.3`, we’re lumping the main effect of therapy and the interaction term together. Running this test in R is straightforward: we just input both models to the `anova()` function, and it will run the exact F -test that I outlined above.

```
> anova( model.1, model.3 )
Analysis of Variance Table

Model 1: mood.gain ~ drug
Model 2: mood.gain ~ drug * therapy
  Res.Df   RSS Df Sum of Sq    F Pr(>F)
1      15 1.392
2      12 0.653  3     0.738 4.52  0.024 *
```

Let’s see if we can reproduce this F -test ourselves. Firstly, if you go back and look at the ANOVA tables that we printed out for `model.1` and `model.3` you can reassure yourself that the RSS values printed in this table really do correspond to the residual sum of squares associated with these two models. So let’s type them in as variables:

```
> ss.res.null <- 1.392
> ss.res.full <- 0.653
```

Now, following the procedure that I described above, we will say that the “between model” sum of squares, is the difference between these two residual sum of squares values. So, if we do the subtraction, we discover that the sum of squares associated with those terms that appear in the full model but not the null model is:

```
> ss.diff <- ss.res.null - ss.res.full
> ss.diff
[1] 0.739
```

Right. Next, as always we need to convert these SS values into MS (mean square) values, which we do by dividing by the degrees of freedom. The degrees of freedom associated with the full-model residuals hasn’t changed from our original ANOVA for `model.3`: it’s the total sample size N , minus the total number of groups G that are relevant to the model. We have 18 people in the trial and 6 possible groups (i.e., 2 therapies \times 3 drugs), so the degrees of freedom here is 12. The degrees of freedom for the null model are calculated similarly. The only difference here is that there are only 3 relevant groups (i.e., 3 drugs), so

the degrees of freedom here is 15. And, because the degrees of freedom associated with the difference is equal to the difference in the two degrees of freedom, we arrive at the conclusion that we have $15 - 12 = 3$ degrees of freedom. Now that we know the degrees of freedom, we can calculate our MS values:

```
> ms.res <- ss.res.full / 12
> ms.diff <- ss.diff / 3
```

Okay, now that we have our two MS values, we can divide one by the other, and obtain an F -statistic ...

```
> F.stat <- ms.diff / ms.res
> F.stat
[1] 4.5268
```

... and, just as we had hoped, this turns out to be identical to the F -statistic that the `anova()` function produced earlier.

16.6

ANOVA as a linear model

One of the most important things to understand about ANOVA and regression is that they're basically the same thing. On the surface of it, you wouldn't think that this is true: after all, the way that I've described them so far suggests that ANOVA is primarily concerned with testing for group differences, and regression is primarily concerned with understanding the correlations between variables. And as far as it goes, that's perfectly true. But when you look under the hood, so to speak, the underlying mechanics of ANOVA and regression are awfully similar. In fact, if you think about it, you've already seen evidence of this. ANOVA and regression both rely heavily on sums of squares (SS), both make use of F tests, and so on. Looking back, it's hard to escape the feeling that Chapters 14 and 15 were a bit repetitive.

The reason for this is that ANOVA and regression are both kinds of **linear models**. In the case of regression, this is kind of obvious. The regression equation that we use to define the relationship between predictors and outcomes *is* the equation for a straight line, so it's quite obviously a linear model. And if that wasn't a big enough clue, the simple fact that the command to run a regression is `lm()` is kind of a hint too. When we use an R formula like `outcome ~ predictor1 + predictor2` what we're really working with is the somewhat uglier linear model:

$$Y_p = b_1 X_{1p} + b_2 X_{2p} + b_0 + \epsilon_p$$

where Y_p is the outcome value for the p -th observation (e.g., p -th person), X_{1p} is the value of the first predictor for the p -th observation, X_{2p} is the value of the second predictor for the p -th observation, the b_1 , b_2 and b_0 terms are our regression coefficients, and ϵ_p is the p -th residual. If we ignore the residuals ϵ_p and just focus on the regression line itself, we get the following formula:

$$\hat{Y}_p = b_1 X_{1p} + b_2 X_{2p} + b_0$$

where \hat{Y}_p is the value of Y that the regression line predicts for person p , as opposed to the actually-observed value Y_p . The thing that isn't immediately obvious is that we can write ANOVA as a linear model as well. However, it's actually pretty straightforward to do this. Let's start with a really simple example: rewriting a 2×2 factorial ANOVA as a linear model.

16.6.1 Some data

To make things concrete, let's suppose that our outcome variable is the `grade` that a student receives in my class, a ratio-scale variable corresponding to a mark from 0% to 100%. There are two predictor variables of interest: whether or not the student turned up to lectures (the `attend` variable), and whether or not the student actually read the textbook (the `reading` variable). We'll say that `attend = 1` if the student attended class, and `attend = 0` if they did not. Similarly, we'll say that `reading = 1` if the student read the textbook, and `reading = 0` if they did not.

Okay, so far that's simple enough. The next thing we need to do is to wrap some maths around this (sorry!). For the purposes of this example, let Y_p denote the `grade` of the p -th student in the class. This is not quite the same notation that we used earlier in this chapter: previously, we've used the notation Y_{rci} to refer to the i -th person in the r -th group for predictor 1 (the row factor) and the c -th group for predictor 2 (the column factor). This extended notation was really handy for describing how the SS values are calculated, but it's a pain in the current context, so I'll switch notation here. Now, the Y_p notation is visually simpler than Y_{rci} , but it has the shortcoming that it doesn't actually keep track of the group memberships! That is, if I told you that $Y_{0,0,3} = 35$, you'd immediately know that we're talking about a student (the 3rd such student, in fact) who didn't attend the lectures (i.e., `attend = 0`) and didn't read the textbook (i.e. `reading = 0`), and who ended up failing the class (`grade = 35`). But if I tell you that $Y_p = 35$ all you know is that the p -th student didn't get a good grade. We've lost some key information here. Of course, it doesn't take a lot of thought to figure out how to fix this: what we'll do instead is introduce two new variables X_{1p} and X_{2p} that keep track of this information. In the case of our hypothetical student, we know that $X_{1p} = 0$ (i.e., `attend = 0`) and $X_{2p} = 0$ (i.e., `reading = 0`). So the data might look like this:

person, p	grade, Y_p	attendance, X_{1p}	reading, X_{2p}
1	90	1	1
2	87	1	1
3	75	0	1
4	60	1	0
5	35	0	0
6	50	0	0
7	65	1	0
8	70	0	1

This isn't anything particularly special, of course: it's exactly the format in which we expect to see our data! In other words, if your data have been stored as a data frame in R then you're probably expecting to see something that looks like the `rtfm.1` data frame:

```
> rtfm.1
  grade attend reading
1    90      1      1
2    87      1      1
3    75      0      1
4    60      1      0
5    35      0      0
6    50      0      0
7    65      1      0
8    70      0      1
```

Well, sort of. I suspect that a few readers are probably frowning a little at this point. Earlier on in the book I emphasised the importance of converting nominal scale variables such as `attend` and `reading` to factors, rather than encoding them as numeric variables. The `rtfm.1` data frame doesn't do this, but the `rtfm.2` data frame does, and so you might instead be expecting to see data like this:

```
> rtfm.2
  grade attend reading
1    90     yes     yes
2    87     yes     yes
3    75      no     yes
4    60     yes      no
5    35      no      no
6    50      no      no
7    65     yes      no
8    70      no     yes
```

However, for the purposes of this section it's important that we be able to switch back and forth between these two different ways of thinking about the data. After all, our goal in this section is to look at some of the mathematics that underpins ANOVA, and if we want to do that we need to be able to see the numerical representation of the data (in `rtfm.1`) as well as the more meaningful factor representation (in `rtfm.2`). In any case, we can use the `xtabs()` function to confirm that this data set corresponds to a balanced design

```
> xtabs( ~ attend + reading, rtfm.2 )
      reading
attend no yes
  no    2    2
  yes   2    2
```

For each possible combination of the `attend` and `reading` variables, we have exactly two students. If we're interested in calculating the mean `grade` for each of these cells, we can use the `aggregate()` function:

```
> aggregate( grade ~ attend + reading, rtfm.2, mean )
  attend reading grade
1      no      no  42.5
2     yes      no  62.5
3      no     yes  72.5
4     yes     yes  88.5
```

Looking at this table, one gets the strong impression that reading the text and attending the class both matter a lot.

16.6.2 ANOVA with binary factors as a regression model

Okay, let's get back to talking about the mathematics. We now have our data expressed in terms of three numeric variables: the continuous variable Y , and the two binary variables X_1 and X_2 . What I want to you to recognise is that our 2×2 factorial ANOVA is *exactly* equivalent to the regression model

$$Y_p = b_1 X_{1p} + b_2 X_{2p} + b_0 + \epsilon_p$$

This is, of course, the exact same equation that I used earlier to describe a two-predictor regression model! The only difference is that X_1 and X_2 are now *binary* variables (i.e., values can only be 0 or 1), whereas in a regression analysis we expect that X_1 and X_2 will be continuous. There's a couple of ways I could try to convince you of this. One possibility would be to do a lengthy mathematical exercise, proving that the two are identical. However, I'm going to go out on a limb and guess that most of the readership of this book will find that to be annoying rather than helpful. Instead, I'll explain the basic ideas, and then rely on R to show that that ANOVA analyses and regression analyses aren't just similar, they're identical

for all intents and purposes.⁹ Let's start by running this as an ANOVA. To do this, we'll use the `rtfm.2` data frame, since that's the one in which I did the proper thing of coding `attend` and `reading` as factors, and I'll use the `aov()` function to do the analysis. Here's what we get...

```
> anova.model <- aov( grade ~ attend + reading, data = rtfm.2 )
> summary( anova.model )
   Df Sum Sq Mean Sq F value    Pr(>F)
attend      1     648     648  21.60 0.00559 ***
reading     1   1568   1568  52.27 0.00079 ***
Residuals   5     150      30
```

So, by reading the key numbers off the ANOVA table and the table of means that we presented earlier, we can see that the students obtained a higher grade if they attended class ($F_{1,5} = 26.1, p = .0056$) and if they read the textbook ($F_{1,5} = 52.3, p = .0008$). Let's make a note of those p -values and those F statistics.

```
> library(effects)
> Effect( c("attend","reading"), anova.model )

attend*reading effect
  reading
attend  no yes
  no  43.5 71.5
  yes 61.5 89.5
```

Now let's think about the same analysis from a linear regression perspective. In the `rtfm.1` data set, we have encoded `attend` and `reading` as if they were numeric predictors. In this case, this is perfectly acceptable. There really is a sense in which a student who turns up to class (i.e. `attend = 1`) has in fact done "more attendance" than a student who does not (i.e. `attend = 0`). So it's not at all unreasonable to include it as a predictor in a regression model. It's a little unusual, because the predictor only takes on two possible values, but it doesn't violate any of the assumptions of linear regression. And it's easy to interpret. If the regression coefficient for `attend` is greater than 0, it means that students that attend lectures get higher grades; if it's less than zero, then students attending lectures get lower grades. The same is true for our `reading` variable.

Wait a second... *why* is this true? It's something that is intuitively obvious to everyone who has taken a few stats classes and is comfortable with the maths, but it *isn't* clear to everyone else at first pass. To see why this is true, it helps to look closely at a few specific students. Let's start by considering the 6th and 7th students in our data set (i.e. $p = 6$ and $p = 7$). Neither one has read the textbook, so in both cases we can set `reading = 0`. Or, to say the same thing in our mathematical notation, we observe $X_{2,6} = 0$ and $X_{2,7} = 0$. However, student number 7 did turn up to lectures (i.e., `attend = 1`, $X_{1,7} = 1$) whereas student number 6 did not (i.e., `attend = 0`, $X_{1,6} = 0$). Now let's look at what happens when we insert these numbers into the general formula for our regression line. For student number 6, the regression predicts that

$$\begin{aligned}\hat{Y}_6 &= b_1 X_{1,6} + b_2 X_{2,6} + b_0 \\ &= (b_1 \times 0) + (b_2 \times 0) + b_0 \\ &= b_0\end{aligned}$$

So we're expecting that this student will obtain a grade corresponding to the value of the intercept term b_0 . What about student 7? This time, when we insert the numbers into the formula for the regression

⁹This is cheating in some respects: because ANOVA and regression are provably the same thing, R is lazy: if you read the help documentation closely, you'll notice that the `aov()` function is actually just the `lm()` function in disguise! But we shan't let such things get in the way of our story, shall we?

line, we obtain the following:

$$\begin{aligned}\hat{Y}_7 &= b_1 X_{1,7} + b_2 X_{2,7} + b_0 \\ &= (b_1 \times 1) + (b_2 \times 0) + b_0 \\ &= b_1 + b_0\end{aligned}$$

Because this student attended class, the predicted grade is equal to the intercept term b_0 plus the coefficient associated with the `attend` variable, b_1 . So, if b_1 is greater than zero, we're expecting that the students who turn up to lectures will get higher grades than those students who don't. If this coefficient is negative, we're expecting the opposite: students who turn up at class end up performing much worse. In fact, we can push this a little bit further. What about student number 1, who turned up to class ($X_{1,1} = 1$) and read the textbook ($X_{2,1} = 1$)? If we plug these numbers into the regression, we get

$$\begin{aligned}\hat{Y}_1 &= b_1 X_{1,1} + b_2 X_{2,1} + b_0 \\ &= (b_1 \times 1) + (b_2 \times 1) + b_0 \\ &= b_1 + b_2 + b_0\end{aligned}$$

So if we assume that attending class helps you get a good grade (i.e., $b_1 > 0$) and if we assume that reading the textbook also helps you get a good grade (i.e., $b_2 > 0$), then our expectation is that student 1 will get a grade that is higher than student 6 and student 7.

And at this point, you won't be at all surprised to learn that the regression model predicts that student 3, who read the book but didn't attend lectures, will obtain a grade of $b_2 + b_0$. I won't bore you with yet another regression formula. Instead, what I'll do is show you the following table of *expected grades*:

		read textbook?	
		no	yes
attended?	no	b_0	$b_0 + b_2$
	yes	$b_0 + b_1$	$b_0 + b_1 + b_2$

As you can see, the intercept term b_0 acts like a kind of "baseline" grade that you would expect from those students who don't take the time to attend class or read the textbook. Similarly, b_1 represents the boost that you're expected to get if you come to class, and b_2 represents the boost that comes from reading the textbook. In fact, if this were an ANOVA you might very well want to characterise b_1 as the main effect of attendance, and b_2 as the main effect of reading! In fact, for a simple 2×2 ANOVA that's *exactly* how it plays out.

Okay, now that we're really starting to see why ANOVA and regression are basically the same thing, let's actually run our regression using the `rtfm.1` data and the `lm()` function to convince ourselves that this is really true. Running the regression in the usual way gives us the following output:¹⁰

```
> regression.model <- lm( grade ~ attend + reading, data = rtfm.1 )
> summary( regression.model )
BLAH BLAH BLAH

Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 43.500     3.354 12.969 4.86e-05 ***
attend       18.000     3.873  4.648  0.00559 **

```

¹⁰In the example given above, I've typed `summary(regression.model)` to get the hypothesis tests. However, the `summary()` function does produce a lot of output, which is why I've used the `BLAH BLAH BLAH` text to hide the unnecessary parts of the output. But in fact, you can use the `coef()` function to do the same job. If you the command `coef(summary(regression.model))` you'll get exactly the same output that I've shown above (minus the `BLAH BLAH BLAH`). Compare and contrast this to the output of `coef(regression.model)`.

```

reading      28.000     3.873    7.230  0.00079 ***
BLAH BLAH BLAH

```

There's a few interesting things to note here. Firstly, notice that the intercept term is 43.5, which is close to the "group" mean of 42.5 observed for those two students who didn't read the text or attend class. Moreover, it's *identical* to the predicted group mean that we pulled out of our ANOVA using the `Effects()` function! Secondly, notice that we have the regression coefficient of $b_1 = 18.0$ for the attendance variable, suggesting that those students that attended class scored 18% higher than those who didn't. So our expectation would be that those students who turned up to class but didn't read the textbook would obtain a grade of $b_0 + b_1$, which is equal to $43.5 + 18.0 = 61.5$. Again, this is similar to the observed group mean of 62.5, and identical to the expected group mean that we pulled from our ANOVA. You can verify for yourself that the same thing happens when we look at the students that read the textbook.

Actually, we can push a little further in establishing the equivalence of our ANOVA and our regression. Look at the *p*-values associated with the `attend` variable and the `reading` variable in the regression output. They're identical to the ones we encountered earlier when running the ANOVA. This might seem a little surprising, since the test used when running our regression model calculates a *t*-statistic and the ANOVA calculates an *F*-statistic. However, if you can remember all the way back to Chapter 9, I mentioned that there's a relationship between the *t*-distribution and the *F*-distribution: if you have some quantity that is distributed according to a *t*-distribution with k degrees of freedom and you square it, then this new squared quantity follows an *F*-distribution whose degrees of freedom are 1 and k . We can check this with respect to the *t* statistics in our regression model. For the `attend` variable we get a *t* value of 4.648. If we square this number we end up with 21.604, which is identical to the corresponding *F* statistic in our ANOVA.

Finally, one last thing you should know. Because R understands the fact that ANOVA and regression are both examples of linear models, it lets you extract the classic ANOVA table from your regression model using the `anova()` function. All you have to do is this:

```

> anova( regression.model )
Analysis of Variance Table

Response: grade
  Df Sum Sq Mean Sq F value    Pr(>F)
attend   1    648     648  21.600 0.0055943 **
reading   1   1568    1568  52.267 0.0007899 ***
Residuals 5    150      30
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1

```

16.6.3 Changing the baseline category

At this point, you're probably convinced that the ANOVA and the regression are actually identical to each other. So there's one last thing I should show you. What happens if I use the data from `rtfm.2` to run the regression? In `rtfm.2`, we coded the `attend` and `reading` variables as factors rather than as numeric variables. Does this matter? It turns out that it doesn't. The only differences are superficial:

```

> regression.model.2 <- lm( grade ~ attend + reading, data = rtfm.2 )
> summary( regression.model.2 )
BLAH BLAH BLAH

Coefficients:

```

```

Estimate Std. Error t value Pr(>|t|)
(Intercept) 43.500     3.354   12.969 4.86e-05 ***
attendyes   18.000     3.873    4.648  0.00559 **
readingyes  28.000     3.873    7.230  0.00079 ***

```

BLAH BLAH BLAH

The only thing that is different is that R labels the two variables differently: the output now refers to `attendyes` and `readingyes`. You can probably guess what this means. When R refers to `readingyes` it's trying to indicate that it is assuming that "yes = 1" and "no = 0". This is important. Suppose we wanted to say that "yes = 0" and "no = 1". We could still run this as a regression model, but now all of our coefficients will go in the opposite direction, because the effect of `readingno` would be referring to the consequences of *not* reading the textbook. To show you how this works, we can use the `relevel()` function in R to change which level of the `reading` variable is set to "0". Here's how it works. First, let's get R to print out the `reading` factor as it currently stands:

```

> rtfm.2$reading
[1] yes yes yes no  no  no  no  yes
Levels: no yes

```

Notice that order in which R prints out the levels is "no" and then "yes". Now let's apply the `relevel()` function:

```

> relevel( x = rtfm.2$reading, ref = "yes" )
[1] yes yes yes no  no  no  no  yes
Levels: yes no

```

R now lists "yes" before "no". This means that R will now treat "yes" as the "reference" level (sometimes called the baseline level) when you include it in an ANOVA. So let's now create a new data frame with our factors recoded...

```

> rtfm.3 <- rtfm.2                                     # copy the old data frame
> rtfm.3$reading <- relevel( rtfm.2$reading, ref="yes" ) # re-level the reading factor
> rtfm.3$attend <- relevel( rtfm.2$attend, ref="yes" )   # re-level the attend factor

```

Finally, let's re-run our regression, this time using the re-coded data:

```

> regression.model.3 <- lm( grade ~ attend + reading, data = rtfm.3 )
> summary( regression.model.3 )
BLAH BLAH BLAH

```

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	89.500	3.354	26.684	1.38e-06 ***
attendno	-18.000	3.873	-4.648	0.00559 **
readingno	-28.000	3.873	-7.230	0.00079 ***

BLAH BLAH BLAH

As you can see, there are now a few changes. Most obviously, the `attendno` and `readingno` effects are both negative, though they're the same magnitude as before: if you *don't* read the textbook, for instance, you should expect your grade to drop by 28% relative to someone who did. The *t*-statistics have reversed sign too. The *p*-values remain the same, of course. The intercept has changed too. In our original regression, the baseline corresponded to a student who didn't attend class and didn't read the textbook, so we got a

value of 43.5 as the expected baseline grade. However, now that we've recoded our variables, the baseline corresponds to a student who has read the textbook and did attend class, and for that student we would expect a grade of 89.5.

16.6.4 How to encode non binary factors as contrasts

At this point, I've shown you how we can view a 2×2 ANOVA into a linear model. And it's pretty easy to see how this generalises to a $2 \times 2 \times 2$ ANOVA or a $2 \times 2 \times 2 \times 2$ ANOVA... it's the same thing, really: you just add a new binary variable for each of your factors. Where it begins to get trickier is when we consider factors that have more than two levels. Consider, for instance, the 3×2 ANOVA that we ran earlier in this chapter using the `clin.trial` data. How can we convert the three-level `drug` factor into a numerical form that is appropriate for a regression?

The answer to this question is pretty simple, actually. All we have to do is realise that a three-level factor can be redescribed as *two* binary variables. Suppose, for instance, I were to create a new binary variable called `druganxitfree`. Whenever the `drug` variable is equal to "`anxitfree`" we set `druganxitfree = 1`. Otherwise, we set `druganxitfree = 0`. This variable sets up a **contrast**, in this case between `anxitfree` and the other two drugs. By itself, of course, the `druganxitfree` contrast isn't enough to fully capture all of the information in our `drug` variable. We need a second contrast, one that allows us to distinguish between `joyzepam` and the placebo. To do this, we can create a second binary contrast, called `drugjoyzepam`, which equals 1 if the drug is `joyzepam`, and 0 if it is not. Taken together, these two contrasts allows us to perfectly discriminate between all three possible drugs. The table below illustrates this:

<code>drug</code>	<code>druganxitfree</code>	<code>drugjoyzepam</code>
"placebo"	0	0
" <code>anxitfree</code> "	1	0
" <code>joyzepam</code> "	0	1

If the drug administered to a patient is a placebo, then both of the two contrast variables will equal 0. If the drug is `Anxitfree`, then the `druganxitfree` variable will equal 1, and `drugjoyzepam` will be 0. The reverse is true for `Joyzepam`: `drugjoyzepam` is 1, and `druganxitfree` is 0.

Creating contrast variables manually is not too difficult to do using basic R commands. For example, here's how we would create the `druganxitfree` variable:

```
> druganxitfree <- as.numeric( clin.trial$drug == "anxitfree" )
> druganxitfree
[1] 0 0 0 1 1 1 0 0 0 0 0 0 1 1 1 0 0 0
```

The `clin.trial$drug == "anxitfree"` part of the command returns a logical vector that has a value of `TRUE` if the drug is `Anxitfree`, and a value of `FALSE` if the drug is `Joyzepam` or the placebo. The `as.numeric()` function just converts `TRUE` to 1 and `FALSE` to 0. Obviously, this command creates the `druganxitfree` variable inside the workspace. If you wanted to add it to the `clin.trial` data frame, you'd use a commmand like this instead:

```
> clin.trial$druganxitfree <- as.numeric( clin.trial$drug == "anxitfree" )
```

You could then repeat this for the other contrasts that you wanted to use. However, it's kind of tedious to do this over and over again for every single contrast that you want to create. To make it a little easier, the `lso` package contains a simple function called `expandFactors()` that will convert every factor in a data frame into a set of contrast variables.¹¹ We can use it to create a new data frame, `clin.trial.2`

¹¹Advanced users may want to look into the `model.matrix()` function, which produces similar output. Alternatively, you can use a command like `contr.treatment(3)[clin.trial$drug,]`. I'll talk about the `contr.treatment()` function later.

that contains the same data as `clin.trial`, but with the two factors represented in terms of the contrast variables:

```
> clin.trial.2 <- expandFactors( clin.trial )
> clin.trial.2
  druganxitfree drugjoyzepam therapyCBT mood.gain
1             0            0            0        0.5
2             0            0            0        0.3
3             0            0            0        0.1
4             1            0            0        0.6
5             1            0            0        0.4
6             1            0            0        0.2
7             0            1            0        1.4
8             0            1            0        1.7

BLAH BLAH BLAH
```

It's not as pretty as the original `clin.trial` data, but it's definitely saying the same thing. We have now recoded our three-level factor in terms of two binary variables, and we've already seen that ANOVA and regression behave the same way for binary variables. However, there are some additional complexities that arise in this case, which we'll discuss in the next section.

16.6.5 The equivalence between ANOVA and regression for non-binary factors

Now we have two different versions of the same data set: our original data frame `clin.trial` in which the `drug` variable is expressed as a single three-level factor, and the expanded data set `clin.trial.2` in which it is expanded into two binary contrasts. Once again, the thing that we want to demonstrate is that our original 3×2 factorial ANOVA is equivalent to a regression model applied to the contrast variables. Let's start by re-running the ANOVA:

```
> drug.anova <- aov( mood.gain ~ drug + therapy, clin.trial )
> summary( drug.anova )
    Df Sum Sq Mean Sq F value Pr(>F)
drug      2   3.45   1.727   26.15 1.9e-05 ***
therapy   1   0.47   0.467    7.08   0.019 *
Residuals 14   0.92   0.066
```

Obviously, there's no surprises here. That's the exact same ANOVA that we ran earlier, except for the fact that I've arbitrarily decided to rename the output variable as `drug.anova` for some stupid reason.¹² Next, let's run a regression, using `draganxitfree`, `drugjoyzepam` and `therapyCBT` as the predictors. Here's what we get:

```
> drug.regression <- lm( mood.gain ~ draganxitfree + drugjoyzepam + therapyCBT, clin.trial.2 )
> summary( drug.regression )

BLAH BLAH BLAH

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  0.289     0.121    2.38    0.032 *
```

¹²Future versions of this book will try to be a bit more consistent with the naming scheme for variables. One of the many problems with having to write a lengthy text very quickly to meet a teaching deadline is that you lose some internal consistency.

druganxitfree	0.267	0.148	1.80	0.094	.
drugjoyzepam	1.033	0.148	6.97	6.6e-06	***
therapyCBT	0.322	0.121	2.66	0.019	*

BLAH BLAH BLAH

Hm. This isn't the same output that we got last time. Not surprisingly, the regression output prints out the results for each of the three predictors separately, just like it did every other time we used `lm()`. On the one hand, we can see that the *p*-value for the `therapyCBT` variable is exactly the same as the one for the `therapy` factor in our original ANOVA, so we can be reassured that the regression model is doing the same thing as the ANOVA did. On the other hand, this regression model is testing the `druganxitfree` contrast and the `drugjoyzepam` contrast *separately*, as if they were two completely unrelated variables. It's not surprising of course, because the poor `lm()` function has no way of knowing that `drugjoyzepam` and `druganxitfree` are actually the two different contrasts that we used to encode our three-level `drug` factor. As far as it knows, `drugjoyzepam` and `druganxitfree` are no more related to one another than `drugjoyzepam` and `therapyCBT`. However, you and I know better. At this stage we're not at all interested in determining whether these two contrasts are individually significant. We just want to know if there's an "overall" effect of drug. That is, what we want R to do is to run some kind of "omnibus" test, one in which the two "drug-related" contrasts are lumped together for the purpose of the test. Sound familiar? This is *exactly* the situation that we discussed in Section 16.5, and it is precisely this situation that the *F*-test is built to handle. All we need to do is specify our null model, which in this case would include the `therapyCBT` predictor, and omit both of the drug-related variables, and then run it through the `anova()` function:

```
> nodrug.regression <- lm( mood.gain ~ therapyCBT, clin.trial.2 )
> anova( nodrug.regression, drug.regression )
Analysis of Variance Table

Model 1: mood.gain ~ therapyCBT
Model 2: mood.gain ~ druganxitfree + drugjoyzepam + therapyCBT
  Res.Df   RSS Df Sum of Sq    F Pr(>F)
1     16 4.38
2     14 0.92  2      3.45 26.1 1.9e-05 ***
```

Ah, that's better. Our *F*-statistic is 26.1, the degrees of freedom are 2 and 14, and the *p*-value is 0.000019. The numbers are identical to the ones we obtained for the main effect of `drug` in our original ANOVA. Once again, we see that ANOVA and regression are essentially the same: they are both linear models, and the underlying statistical machinery for ANOVA is identical to the machinery used in regression. The importance of this fact should not be understated. Throughout the rest of this chapter we're going to rely heavily on this idea.

16.6.6 Degrees of freedom as parameter counting!

At long last, I can finally give a definition of degrees of freedom that I am happy with. Degrees of freedom are defined in terms of the number of parameters that have to be estimated in a model. For a regression model or an ANOVA, the number of parameters corresponds to the number of regression coefficients (i.e. *b*-values), including the intercept. Keeping in mind that any *F*-test is always a comparison between two models, the first *df* is the difference in the number of parameters. For example, model comparison above, the null model (`mood.gain ~ therapyCBT`) has two parameters: there's one regression coefficient for the `therapyCBT` variable, and a second one for the intercept. The alternative model (`mood.gain ~ druganxitfree + drugjoyzepam + therapyCBT`) has four parameters: one regression coefficient for each of the three contrasts, and one more for the intercept. So the degrees of freedom associated with the *difference* between these two models is $df_1 = 4 - 2 = 2$.

What about the case when there doesn't seem to *be* a null model? For instance, you might be thinking of the F -test that appears at the very bottom of the regression output. I originally described that as a test of the regression model as a whole. However, that is still a comparison between two models. The null model is the trivial model that only includes an intercept, which is written as `outcome ~ 1` in R, and the alternative model is the full regression model. The null model in this case contains 1 regression coefficient, for the intercept term. The alternative model contains $K + 1$ regression coefficients, one for each of the K predictor variables and one more for the intercept. So the df value that you see in this F test is equal to $df_1 = K + 1 - 1 = K$.

What about the second df value that appears in the F -test? This always refers to the degrees of freedom associated with the residuals. It is possible to think of this in terms of parameters too, but in a slightly counterintuitive way. Think of it like this: suppose that the total number of observations across the study as a whole is N . If you wanted to *perfectly* describe each of these N values, you need to do so using, well... N numbers. When you build a regression model, what you're really doing is specifying some of the numbers need to perfectly describe the data. If your model has K predictors and an intercept, then you've specified $K + 1$ numbers. So, without bothering to figure out exactly *how* this would be done, how many *more* numbers do you think are going to be needed to transform a $K + 1$ parameter regression model into a perfect redescription of the raw data? If you found yourself thinking that $(K + 1) + (N - K - 1) = N$, and so the answer would have to be $N - K - 1$, well done! That's exactly right: in principle you can imagine an absurdly complicated regression model that includes a parameter for every single data point, and it would of course provide a perfect description of the data. This model would contain N parameters in total, but we're interested in the difference between the number of parameters required to describe this full model (i.e. N) and the number of parameters used by the simpler regression model that you're actually interested in (i.e., $K + 1$), and so the second degrees of freedom in the F test is $df_2 = N - K - 1$, where K is the number of predictors (in a regression model) or the number of contrasts (in an ANOVA). In the example I gave above, there are $N = 18$ observations in the data set, and $K + 1 = 4$ regression coefficients associated with the ANOVA model, so the degrees of freedom for the residuals is $df_2 = 18 - 4 = 14$.

16.6.7 A postscript

There's one last thing I want to mention in this section. In the previous example, I used the `aov()` function to run an ANOVA using the `clin.trial` data which codes `drug` variable as a single factor. I also used the `lm()` function to run a regression using the `clin.trial` data in which we have two separate contrasts describing the drug. However, it's also possible to use the `lm()` function on the the original data. That is, you could use a command like this:

```
> drug.lm <- lm( mood.gain ~ drug + therapy, clin.trial )
```

The fact that `drug` is a three-level factor does not matter. As long as the `drug` variable has been declared to be a factor, R will automatically translate it into two binary contrast variables, and will perform the appropriate analysis. After all, as I've been saying throughout this section, ANOVA and regression are both linear models, and `lm()` is the function that handles linear models. In fact, the `aov()` function doesn't actually do very much of the work when you run an ANOVA using it: internally, R just passes all the hard work straight to `lm()`. However, I want to emphasise again that it is critical that your factor variables are declared as such. If `drug` were declared to be a numeric variable, then R would be happy to treat it as one. After all, it might be that `drug` refers to the number of drugs that one has taken in the past, or something that is genuinely numeric. R won't second guess you here. It assumes your factors are factors and your numbers are numbers. Don't make the mistake of encoding your factors as numbers, or R will run the wrong analysis. This is *not* a flaw in R: it is *your* responsibility as the analyst to make sure you're specifying the right model for your data. Software really can't be trusted with this sort of thing.

Okay, warnings aside, it's actually kind of neat to run your ANOVA using the `lm()` function in the

way I did above. Because you've called the `lm()` function, the `summary()` that R pulls out is formatted like a regression. To save space I won't show you the output here, but you can easily verify this by typing

```
> summary( drug.lm )
```

However, because the `drug` and `therapy` variables were both factors, the `anova()` function actually knows which contrasts to group together for the purposes of running the F -tests, so you can extract the classic ANOVA table. Again, I won't reproduce the output here since it's identical to the ANOVA table I showed at the start of the section, but it's worth trying the following command

```
> anova( drug.lm )
```

just to see for yourself. However, this behaviour of the `anova()` function only occurs when the predictor variables are factors. If we try a command like `anova(drug.regression)`, the output will continue to treat `druganxitfree` and `drugjoyzepam` as if they were two distinct binary factors. This is because in the `drug.regression` model we included all the contrasts as "raw" variables, so R had no idea which ones belonged together. However, when we ran the `drug.lm` model, we gave R the original factor variables, so it does know which contrasts go together. The behaviour of the `anova()` function reflects that.

16.7

Different ways to specify contrasts

In the previous section, I showed you a method for converting a factor into a collection of contrasts. In the method I showed you, we specify a set of binary variables, in which define a table like this one:

drug	druganxitfree	drugjoyzepam
"placebo"	0	0
"anxitfree"	1	0
"joyzepam"	0	1

Each row in the table corresponds to one of the factor levels, and each column corresponds to one of the contrasts. This table, which always has one more row than columns, has a special name: it is called a **contrast matrix**. However, there are lots of different ways to specify a contrast matrix. In this section I discuss a few of the standard contrast matrices that statisticians use, and how you can use them in R. If you're planning to read the section on unbalanced ANOVA later on (Section 16.10) it's worth reading this section carefully. If not, you can get away with skimming it, because the choice of contrasts doesn't matter much for balanced designs.

16.7.1 Treatment contrasts

In the particular kind of contrasts that I've described above, one level of the factor is special, and acts as a kind of "baseline" category (i.e., `placebo` in our example), against which the other two are defined. The name for these kinds of contrast is **treatment contrasts**. The name reflects the fact that these contrasts are quite natural and sensible when one of the categories in your factor really is special because it actually does represent a baseline. That makes sense in our clinical trial example: the `placebo` condition corresponds to the situation where you don't give people any real drugs, and so it's special. The other two conditions are defined in relation to the placebo: in one case you replace the placebo with Anxitfree, and in the other case you replace it with Joyzepam.

R comes with a variety of functions that can generate different kinds of contrast matrices. For example, the table shown above is a matrix of treatment contrasts for a factor that has 3 levels. But suppose I want a matrix of treatment contrasts for a factor with 5 levels? The `contr.treatment()` function will do this:

```
> contr.treatment( n=5 )
  2 3 4 5
1 0 0 0 0
2 1 0 0 0
3 0 1 0 0
4 0 0 1 0
5 0 0 0 1
```

Notice that, by default, the *first* level of the factor is always treated as the baseline category (i.e., it's the one that has all zeros, and doesn't have an explicit contrast associated with it). In Section 16.6.3 I mentioned that you can use the `relevel()` function to change which category is the first level of the factor.¹³ There's also a special function in R called `contr.SAS()` that generates a treatment contrast matrix in which the *last* category is treated as the baseline:

```
> contr.SAS( n=5 )
  1 2 3 4
1 1 0 0 0
2 0 1 0 0
3 0 0 1 0
4 0 0 0 1
5 0 0 0 0
```

However, you can actually select any category you like as the baseline within the `contr.treatment()` function, by specifying the `base` argument in that function. See the help documentation for more details.

16.7.2 Helmert contrasts

Treatment contrasts are useful for a lot of situations, and they're the default in R. However, they make most sense in the situation when there really is a baseline category, and you want to assess all the other groups in relation to that one. In other situations, however, no such baseline category exists, and it may make more sense to compare each group to the mean of the other groups. This is where **Helmert contrasts**, generated by the `contr.helmert()` function, can be useful. The idea behind Helmert contrasts is to compare each group to the mean of the "previous" ones. That is, the first contrast represents the difference between group 2 and group 1, the second contrast represents the difference between group 3 and the mean of groups 1 and 2, and so on. This translates to a contrast matrix that looks like this:

```
> contr.helmert( n=5 )
 [,1] [,2] [,3] [,4]
1   -1   -1   -1   -1
2    1   -1   -1   -1
3    0    2   -1   -1
4    0    0    3   -1
5    0    0    0    4
```

One useful thing about Helmert contrasts is that every contrast sums to zero (i.e., all the columns sum to zero). This has the consequence that, when we interpret the ANOVA as a regression, the intercept term corresponds to the grand mean $\mu_{..}$) if we are using Helmert contrasts. Compare this to treatment

¹³The `lsr` package contains a more general function called `permuteLevels()` that can shuffle them in any way you like.

contrasts, in which the intercept term corresponds to the group mean for the baseline category. This property can be very useful in some situations. It doesn't matter very much if you have a balanced design, which we've been assuming so far, but it will turn out to be important later when we consider unbalanced designs in Section 16.10. In fact, the main reason why I've even bothered to include this section on specifying is that contrasts become important if you want to understand unbalanced ANOVA.

16.7.3 Sum to zero contrasts

The third option that I should briefly mention are “sum to zero” contrasts, which are used to construct pairwise comparisons between groups. Specifically, each contrast encodes the difference between one of the groups and a baseline category, which in this case corresponds to the last group:

```
> contr.sum( n=5 )
 [,1] [,2] [,3] [,4]
 1     1     0     0     0
 2     0     1     0     0
 3     0     0     1     0
 4     0     0     0     1
 5    -1    -1    -1    -1
```

Much like Helmert contrasts, we see that each column sums to zero, which means that the intercept term corresponds to the grand mean when ANOVA is treated as a regression model. When interpreting these contrasts, the thing to recognise is that each of these contrasts is a pairwise comparison between group 5 and one of the other four groups. Specifically, contrast 1 corresponds to a “group 1 minus group 5” comparison, contrast 2 corresponds to a “group 2 minus group 5” comparison, and so on.

16.7.4 Viewing and setting the default contrasts in R

Every factor variable in R is associated with a contrast matrix. It has to be, otherwise R wouldn't be able to run ANOVAs properly! If you don't specify one explicitly, or R will implicitly specify one for you. Here's what I mean. When I created the `clin.trial` data, I didn't specify any contrast matrix for either of the factors. You can see this by using the `attr()` function to print out the “contrasts” attribute of the factors. For example:

```
> attr( clin.trial$drug, "contrasts" )
NULL
```

The `NULL` output here means that R is telling you that the `drug` factor doesn't have any attribute called “contrasts” for which it has any data. There is no contrast matrix stored anywhere explicitly for this factor. However, if we now ask R to tell us what contrasts are set up for this factor, it give us this:

```
> contrasts( clin.trial$drug )
      anxifree joyzepam
placebo        0        0
anxifree       1        0
joyzepam       0        1
```

These are the same treatment contrast that we set up manually in Section 16.6. How did R know to set up treatment contrasts, even though I never actually told it anything about what contrasts I wanted? The answer is that R has a hidden list of default “options” that it looks up to resolve situations like this. You can print out all of the options by typing `options()` at the command prompt, but it's not a very

enlightening read. There are a lot of options, and we're only interested in contrasts right now. Instead of printing out all of the options, we can ask for just one, like this:

```
> options("contrasts")
$contrasts
  unordered          ordered
"contr.treatment"    "contr.poly"
```

What this is telling us is that the default contrasts for unordered factors (i.e., nominal scale variables) are treatment contrasts, and the default for ordered factors (i.e., interval scale variables) are “polynomial” contrasts. I don't discuss ordered factors much in this book, and so I won't go into what polynomial contrasts are all about. The key thing is that the `options()` function also allows you to reset these defaults (though only for the current session: they'll revert to the original settings once you close R). Here's the command:

```
> options(contrasts = c("contr.helmert", "contr.poly"))
```

Once we've done this, we can inspect the contrast settings again:

```
> options("contrasts")
$contrasts
[1] "contr.helmert" "contr.poly"
```

Now we see that the default contrasts for unordered factors have changed. So if I now ask R to tell me what contrasts are associated with the `drug` factor, it gives a different answer because I changed the default:

```
> contrasts( clin.trial$drug )
 [,1] [,2]
 placebo   -1   -1
 anxifree    1   -1
 joyzepam     0    2
```

Those are Helmert contrasts. In general, if you're changing the default settings for something in R, it's a good idea to reset them to their original values once you're done. So let's do that:

```
> options(contrasts = c("contr.treatment", "contr.poly"))
```

16.7.5 Setting the contrasts for a single factor

In the previous section, I showed you how to alter the default contrasts. However, suppose that all you really want to do is change the contrasts associated with a single factor, and leave the defaults as they are. To do this, what you need to do is specifically assign the contrast matrix as an “attribute” of the factor. This is easy to do via the `contrasts()` function. For instance, suppose I wanted to use sum to zero contrasts for the `drug` factor, but keep the default treatment contrasts for everything else. I could do that like so:

```
> contrasts( clin.trial$drug ) <- contr.sum(3)
```

And if I now inspect the contrasts, I get the following

```
> contrasts( clin.trial$drug )
      [,1] [,2]
placebo     1    0
anxifree    0    1
joyzepam   -1   -1
```

However, the contrasts for everything else will still be the defaults. You can check that we have actually made a specific change to the factor itself by checking to see if it now has an attribute, using the command `attr(clin.trial$drug, "contrasts")`. This will print out the same output shown above, because the contrast has in fact been attached to the `drug` factor, and does not rely on the defaults. If you want to wipe the attribute and revert the defaults, use a command like this:

```
> contrasts( clin.trial$drug ) <- NULL
```

16.7.6 Setting the contrasts for a single analysis

One last way of changing contrasts. You might find yourself wanting to change the contrasts only for one specific analysis. That's allowed too, because the `aov()` and `lm()` functions have a `contrasts` argument that you can use. To change contrasts for one specific analysis, we first set up a list variable that names¹⁴ the contrast types that you want to use for each of the factors:

```
> my.contrasts <- list( drug = contr.helmert, therapy = contr.helmert )
```

Next, fit the ANOVA model in the usual way, but this time we'll specify the `contrasts` argument:

```
> mod <- aov( mood.gain ~ drug*therapy, clin.trial, contrasts = my.contrasts )
```

If you try a command like `summary(aov)` you won't see any difference in the output because the choice of contrasts does not affect the outcome when you have a balanced design (this won't always be true later on). However, if you want to check that it has actually worked, you can inspect the value of `mod$contrasts`:

```
> mod$contrasts
$drug
      [,1] [,2]
placebo   -1   -1
anxifree    1   -1
joyzepam    0    2

$therapy
      [,1]
no.therapy -1
CBT         1
```

As you can see, for the purposes of this one particular ANOVA, R has used Helmert contrasts for both variables. If I had omitted the part of the command that specified the `contrasts` argument, you'd be looking at treatment contrasts here because it would have reverted to whatever values the `contrasts()` function prints out for each of the factors.

¹⁴Technically, this list actually stores the functions themselves. R allows lists to contain functions, which is really neat for advanced purposes, but not something that matters for this book.

Post hoc tests

Time to switch to a different topic. Let's suppose you've done your ANOVA, and it turns out that you obtained some significant effects. Because of the fact that the F -tests are "omnibus" tests that only really test the null hypothesis that there are no differences among groups, obtaining a significant effect doesn't tell you which groups are different to which other ones. We discussed this issue back in Chapter 14, and in that chapter our solution was to run t -tests for all possible pairs of groups, making corrections for multiple comparisons (e.g., Bonferroni, Holm) to control the Type I error rate across all comparisons. The methods that we used back in Chapter 14 have the advantage of being relatively simple, and being the kind of tools that you can use in a lot of different situations where you're testing multiple hypotheses, but they're not necessarily the best choices if you're interested in doing efficient post hoc testing in an ANOVA context. There are actually quite a lot of different methods for performing multiple comparisons in the statistics literature (Hsu, 1996), and it would be beyond the scope of an introductory text like this one to discuss all of them in any detail.

That being said, there's one tool that I do want to draw your attention to, namely Tukey's "Honestly Significant Difference", or **Tukey's HSD** for short. For once, I'll spare you the formulas, and just stick to the qualitative ideas. The basic idea in Tukey's HSD is to examine all relevant pairwise comparisons between groups, and it's only really appropriate to use Tukey's HSD if it is *pairwise* differences that you're interested in.¹⁵ For instance, in `model.2`, where we specified a main effect for drug and a main effect of therapy, we would be interested in the following four comparisons:

- The difference in mood gain for people given Anxifree versus people given the placebo.
- The difference in mood gain for people given Joyzepam versus people given the placebo.
- The difference in mood gain for people given Anxifree versus people given Joyzepam.
- The difference in mood gain for people treated with CBT and people given no therapy.

For any one of these comparisons, we're interested in the true difference between (population) group means. Tukey's HSD constructs **simultaneous confidence intervals** for all four of these comparisons. What we mean by 95% "simultaneous" confidence interval is that there is a 95% probability that *all* of these confidence intervals contain the relevant true value. Moreover, we can use these confidence intervals to calculate an adjusted p value for any specific comparison.

The `TukeyHSD()` function in R is pretty easy to use: you simply input the model that you want to run the post hoc tests for. For example, if we were looking to run post hoc tests for `model.2`, here's the command we would use:

```
> TukeyHSD( model.2 )

Tukey multiple comparisons of means
 95% family-wise confidence level

Fit: aov(formula = mood.gain ~ drug + therapy, data = clin.trial)

$drug
```

¹⁵If, for instance, you actually would find yourself interested to know if Group A is significantly different from the mean of Group B and Group C, then you need to use a different tool (e.g., Scheffe's method, which is more conservative, and beyond the scope this book). However, in most cases you probably are interested in pairwise group differences, so Tukey's HSD is a pretty useful thing to know about.

```

          diff      lwr      upr      p adj
anxitfree-placebo 0.2666667 -0.1216321 0.6549655 0.2062942
joyzepam-placebo 1.0333333 0.6450345 1.4216321 0.0000186
joyzepam-anxitfree 0.7666667 0.3783679 1.1549655 0.0003934

$therapy
          diff      lwr      upr      p adj
CBT-no.therapy 0.3222222 0.0624132 0.5820312 0.0186602

```

The output here is (I hope) pretty straightforward. The first comparison, for example, is the Anxitfree versus placebo difference, and the first part of the output indicates that the observed difference in group means is .27. The next two numbers indicate that the 95% (simultaneous) confidence interval for this comparison runs from -0.12 to 0.65 . Because the confidence interval for the difference includes 0, we cannot reject the null hypothesis that the two group means are identical, and so we're not all that surprised to see that the adjusted p -value is .21. In contrast, if you look at the next line, we see that the observed difference between Joyzepam and the placebo is 1.03, and the 95% confidence interval runs from 0.64 to 1.42 . Because the interval excludes 0, we see that the result is significant ($p < .001$).

So far, so good. What about the situation where your model includes interaction terms? For instance, in `model.3` we allowed for the possibility that there is an interaction between drug and therapy. If that's the case, the number of pairwise comparisons that we need to consider starts to increase. As before, we need to consider the three comparisons that are relevant to the main effect of `drug` and the one comparison that is relevant to the main effect of `therapy`. But, if we want to consider the possibility of a significant interaction (and try to find the group differences that underpin that significant interaction), we need to include comparisons such as the following:

- The difference in mood gain for people given Anxitfree and treated with CBT, versus people given the placebo and treated with CBT
- The difference in mood gain for people given Anxitfree and given no therapy, versus people given the placebo and given no therapy.
- etc

There are quite a lot of these comparisons that you need to consider. So, when we run the `TukeyHSD()` command for `model.3` we see that it has made a *lot* of pairwise comparisons (19 in total). Here's the output:

```

> TukeyHSD( model.3 )
Tukey multiple comparisons of means
 95% family-wise confidence level

Fit: aov(formula = mood.gain ~ drug * therapy, data = clin.trial)

$drug
          diff      lwr      upr      p adj
anxitfree-placebo 0.2666667 -0.09273475 0.6260681 0.1597148
joyzepam-placebo 1.0333333 0.67393191 1.3927348 0.0000160
joyzepam-anxitfree 0.7666667 0.40726525 1.1260681 0.0002740

$therapy
          diff      lwr      upr      p adj
CBT-no.therapy 0.3222222 0.08256504 0.5618794 0.012617

$`drug:therapy`

```

	diff	lwr	upr	p adj
anxitfree:no.therapy-placebo:no.therapy	0.1000000	-0.539927728	0.7399277	0.9940083
joyzepam:no.therapy-placebo:no.therapy	1.16666667	0.526738939	1.8065944	0.0005667
placebo:CBT-placebo:no.therapy	0.3000000	-0.339927728	0.9399277	0.6280049
anxitfree:CBT-placebo:no.therapy	0.73333333	0.093405606	1.3732611	0.0218746
joyzepam:CBT-placebo:no.therapy	1.2000000	0.560072272	1.8399277	0.0004380
joyzepam:no.therapy-anxitfree:no.therapy	1.06666667	0.426738939	1.7065944	0.0012553
placebo:CBT-anxitfree:no.therapy	0.2000000	-0.439927728	0.8399277	0.8917157
anxitfree:CBT-anxitfree:no.therapy	0.63333333	-0.006594394	1.2732611	0.0529812
joyzepam:CBT-anxitfree:no.therapy	1.1000000	0.460072272	1.7399277	0.0009595
placebo:CBT-joyzepam:no.therapy	-0.86666667	-1.506594394	-0.2267389	0.0067639
anxitfree:CBT-joyzepam:no.therapy	-0.43333333	-1.073261061	0.2065944	0.2750590
joyzepam:CBT-joyzepam:no.therapy	0.03333333	-0.606594394	0.6732611	0.9999703
anxitfree:CBT-placebo:CBT	0.43333333	-0.206594394	1.0732611	0.2750590
joyzepam:CBT-placebo:CBT	0.9000000	0.260072272	1.5399277	0.0050693
joyzepam:CBT-anxitfree:CBT	0.46666667	-0.173261061	1.1065944	0.2139229

It looks pretty similar to before, but with a lot more comparisons made.

16.9

The method of planned comparisons

Okay, I have a confession to make. I haven't had time to write this section, but I think the method of planned comparisons is important enough to deserve a quick discussion. In our discussions of multiple comparisons, in the previous section and back in Chapter 14, I've been assuming that the tests you want to run are genuinely post hoc. For instance, in our drugs example above, maybe you thought that the drugs would all have different effects on mood (i.e., you hypothesised a main effect of drug), but you didn't have any specific hypothesis about how they would be different, nor did you have any real idea about *which* pairwise comparisons would be worth looking at. If that is the case, then you really have to resort to something like Tukey's HSD to do your pairwise comparisons.

The situation is rather different, however, if you genuinely did have real, specific hypotheses about which comparisons are of interest, and you *never ever* have any intention to look at any other comparisons besides the ones that you specified ahead of time. When this is true, and if you honestly and rigourously stick to your noble intentions to not run any other comparisons (even when the data look like they're showing you deliciously significant effects for stuff you didn't have a hypothesis test for), then it doesn't really make a lot of sense to run something like Tukey's HSD, because it makes corrections for a whole bunch of comparisons that you never cared about and never had any intention of looking at. Under those circumstances, you can safely run a (limited) number of hypothesis tests without making an adjustment for multiple testing. This situation is known as the **method of planned comparisons**, and it is sometimes used in clinical trials. In a later version of this book, I intend to talk a lot more about planned comparisons, but I'm writing to a deadline here, in order to get the book available to serve as lecture notes for the 2013 class, and so I haven't been able to write this yet.

Factorial ANOVA 3: unbalanced designs

Factorial ANOVA is a very handy thing to know about. It's been one of the standard tools used to analyse experimental data for many decades, and you'll find that you can't read more than two or three papers in psychology without running into an ANOVA in there somewhere. However, there's one huge difference between the ANOVAs that you'll see in a lot of real scientific articles and the ANOVA that I've just described: in real life, we're rarely lucky enough to have perfectly balanced designs. For one reason or another, it's typical to end up with more observations in some cells than in others. Or, to put it another way, we have an **unbalanced design**.

Unbalanced designs need to be treated with a lot more care than balanced designs, and the statistical theory that underpins them is a lot messier. It might be a consequence of this messiness, or it might be a shortage of time, but my experience has been that undergraduate research methods classes in psychology have a nasty tendency to ignore this issue completely. A lot of stats textbooks tend to gloss over it too. The net result of this, I think, is that a lot of active researchers in the field don't actually know that there's several different "types" of unbalanced ANOVAs, and they produce quite different answers. In fact, reading the psychological literature, I'm kind of amazed at the fact that most people who report the results of an unbalanced factorial ANOVA don't actually give you enough details to reproduce the analysis: I secretly suspect that most people don't even realise that their statistical software package is making a whole lot of substantive data analysis decisions on their behalf. It's actually a little terrifying, when you think about it. So, if you want to avoid handing control of your data analysis to stupid software, read on...

16.10.1 The coffee data

As usual, it will help us to work with some data. The `coffee.Rdata` file contains a hypothetical data set (the `coffee` data frame) that produces an unbalanced 3×2 ANOVA. Suppose we were interested in finding out whether or not the tendency of people to `babble` when they have too much coffee is purely an effect of the coffee itself, or whether there's some effect of the `milk` and `sugar` that people add to the coffee. Suppose we took 18 people, and gave them some coffee to drink. The amount of coffee / caffeine was held constant, and we varied whether or not milk was added: so `milk` is a binary factor with two levels, `"yes"` and `"no"`. We also varied the kind of sugar involved. The coffee might contain `"real"` sugar, or it might contain `"fake"` sugar (i.e., artificial sweetener), or it might contain `"none"` at all, so the `sugar` variable is a three level factor. Our outcome variable is a continuous variable that presumably refers to some psychologically sensible measure of the extent to which someone is "babbling". The details don't really matter for our purpose. To get a sense of what the data look like, we use the `some()` function in the `car` package. The `some()` function randomly picks a few of the observations in the data frame to print out, which is often very handy:

```
> some( coffee )
   milk sugar babble
 4  yes  real    5.6
 5  yes  real    4.9
 7  yes  none    3.8
 8  yes  none    3.7
12  yes  fake    5.6
14  no   real    6.1
15  no   real    6.3
16  no   none    5.5
```

```

17 no none 5.0
18 yes fake 5.0

```

If we use the `aggregate()` function to quickly produce a table of means, we get a strong impression that there are differences between the groups:

```

> aggregate( babble ~ milk + sugar, coffee, mean )
   milk sugar babble
1 yes  none 3.700
2 no   none 5.550
3 yes  fake 5.800
4 no   fake 4.650
5 yes  real  5.100
6 no   real  5.875

```

This is especially true when we compare these means to the standard deviations for the `babble` variable, which you can calculate using `aggregate()` in much the same way. Across groups, this standard deviation varies from .14 to .71, which is fairly small relative to the differences in group means.¹⁶ So far, it's looking like a straightforward factorial ANOVA, just like we did earlier. The problem arises when we check to see how many observations we have in each group:

```

> xtabs( ~ milk + sugar, coffee )
   sugar
milk  none fake real
yes    3    2    3
no     2    4    4

```

This violates one of our original assumptions, namely that the number of people in each group is the same. We haven't really discussed how to handle this situation.

16.10.2 “Standard ANOVA” does not exist for unbalanced designs

Unbalanced designs lead us to the somewhat unsettling discovery that there isn't really any one thing that we might refer to as a standard ANOVA. In fact, it turns out that there are *three* fundamentally different ways¹⁷ in which you might want to run an ANOVA in an unbalanced design. If you have a balanced design, all three versions produce identical results, with the sums of squares, *F*-values etc all conforming to the formulas that I gave at the start of the chapter. However, when your design is unbalanced they don't give the same answers. Furthermore, they are not all equally appropriate to every situation: some methods will be more appropriate to your situation than others. Given all this, it's important to understand what the different types of ANOVA are and how they differ from one another.

The first kind of ANOVA is conventionally referred to as **Type I sum of squares**. I'm sure you can guess what the other two are called. The “sum of squares” part of the name was introduced by

¹⁶This discrepancy in standard deviations might (and should) make you wonder if we have a violation of the homogeneity of variance assumption. I'll leave it as an exercise for the reader to check this using the `leveneTest()` function.

¹⁷Actually, this is a bit of a lie. ANOVAs can vary in other ways besides the ones I've discussed in this book. For instance, I've completely ignored the difference between fixed-effect models, in which the levels of a factor are “fixed” by the experimenter or the world, and random-effect models, in which the levels are random samples from a larger population of possible levels (this book only covers fixed-effect models). Don't make the mistake of thinking that this book – or any other one – will tell you “everything you need to know” about statistics, any more than a single book could possibly tell you everything you need to know about psychology, physics or philosophy. Life is too complicated for that to *ever* be true. This isn't a cause for despair, though. Most researchers get by with a basic working knowledge of ANOVA that doesn't go any further than this book does. I just want you to keep in mind that this book is only the beginning of a very long story, not the whole story.

the SAS statistical software package, and has become standard nomenclature, but it's a bit misleading in some ways. I think the logic for referring to them as different types of sum of squares is that, when you look at the ANOVA tables that they produce, the key difference in the numbers is the SS values. The degrees of freedom don't change, the MS values are still defined as SS divided by df, etc. However, what the terminology gets wrong is that it hides the reason *why* the SS values are different from one another. To that end, it's a lot more helpful to think of the three different kinds of ANOVA as three different *hypothesis testing strategies*. These different strategies lead to different SS values, to be sure, but it's the strategy that is the important thing here, not the SS values themselves. Recall from Section 16.5 and 16.6 that any particular F -test is best thought of as a comparison between two linear models. So when you're looking at an ANOVA table, it helps to remember that each of those F -tests corresponds to a *pair* of models that are being compared. Of course, this leads naturally to the question of *which* pair of models is being compared. This is the fundamental difference between ANOVA Types I, II and III: each one corresponds to a different way of choosing the model pairs for the tests.

16.10.3 Type I sum of squares

The Type I method is sometimes referred to as the “sequential” sum of squares, because it involves a process of adding terms to the model one at a time. Consider the coffee data, for instance. Suppose we want to run the full 3×2 factorial ANOVA, including interaction terms. The full model, as we've discussed earlier, is expressed by the R formula `babble ~ sugar + milk + sugar:milk`, though we often shorten it by using the `sugar * milk` notation. The Type I strategy builds this model up sequentially, starting from the simplest possible model and gradually adding terms.

The simplest possible model for the data would be one in which neither milk nor sugar is assumed to have any effect on babbling. The only term that would be included in such a model is the intercept, and in R formula notation we would write it as `babble ~ 1`. This is our initial null hypothesis. The next simplest model for the data would be one in which only one of the two main effects is included. In the coffee data, there are two different possible choices here, because we could choose to add milk first or to add sugar first (pardon the pun). The order actually turns out to matter, as we'll see later, but for now let's just make a choice arbitrarily, and pick sugar. So the second model in our sequence of models is `babble ~ sugar`, and it forms the alternative hypothesis for our first test. We now have our first hypothesis test:

Null model: `babble ~ 1`
Alternative model: `babble ~ sugar`

This comparison forms our hypothesis test of the main effect of `sugar`. The next step in our model building exercise is to add the other main effect term, so the next model in our sequence is `babble ~ sugar + milk`. The second hypothesis test is then formed by comparing the following pair of models:

Null model: `babble ~ sugar`
Alternative model: `babble ~ sugar + milk`

This comparison forms our hypothesis test of the main effect of `milk`. In one sense, this approach is very elegant: the alternative hypothesis from the first test forms the null hypothesis for the second one. It is in this sense that the Type I method is strictly sequential. Every test builds directly on the results of the last one. However, in another sense it's very inelegant, because there's a strong asymmetry between the two tests. The test of the main effect of `sugar` (the first test) completely ignores `milk`, whereas the test of the main effect of `milk` (the second test) does take `sugar` into account. In any case, the fourth model in our sequence is now the full model, `babble ~ sugar + milk + sugar:milk`, and the corresponding

hypothesis test is

```
Null model:      babble ~ sugar + milk
Alternative model: babble ~ sugar + milk + sugar:milk
```

Type I sum of squares is the default hypothesis testing method in R, so it's easy to produce the results from a Type I analysis. We just type in the same commands that we always did. Since we've now reached the point that we don't need to hide the fact that ANOVA and regression are both linear models, I'll use the `lm()` function to run the analyses:

```
> mod <- lm( babble ~ sugar + milk + sugar:milk, coffee )
> anova( mod )
Analysis of Variance Table

Response: babble
          Df Sum Sq Mean Sq F value    Pr(>F)
sugar      2 3.5575 1.77876  6.7495 0.010863 *
milk       1 0.9561 0.95611  3.6279 0.081061 .
sugar:milk 2 5.9439 2.97193 11.2769 0.001754 **
Residuals 12 3.1625 0.26354
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

Leaving aside for one moment the question of how this result should be interpreted, let's take note of the fact that our three p -values are .0109, .0811 and .0018 respectively. Next, let's see if we can replicate the analysis using tools that we're a little more familiar with. First, let's fit all four models:

```
> mod.1 <- lm( babble ~ 1, coffee )
> mod.2 <- lm( babble ~ sugar, coffee )
> mod.3 <- lm( babble ~ sugar + milk, coffee )
> mod.4 <- lm( babble ~ sugar + milk + sugar:milk, coffee )
```

To run the first hypothesis test comparing `mod.1` to `mod.2` we can use the command `anova(mod.1, mod.2)` in much the same way that we did in Section 16.5. Similarly, we can use the commands `anova(mod.2, mod.3)` and `anova(mod.3, mod.4)` and to run the second and third hypothesis tests. However, rather than run each of those commands separately, we can enter the full sequence of models like this:

```
> anova( mod.1, mod.2, mod.3, mod.4 )
Analysis of Variance Table

Model 1: babble ~ 1
Model 2: babble ~ sugar
Model 3: babble ~ sugar + milk
Model 4: babble ~ sugar + milk + sugar:milk
  Res.Df   RSS Df Sum of Sq    F    Pr(>F)
1     17 13.6200
2     15 10.0625  2    3.5575  6.7495 0.010863 *
3     14  9.1064  1    0.9561  3.6279 0.081061 .
4     12  3.1625  2    5.9439 11.2769 0.001754 **
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

This output is rather more verbose than the last one, but it's telling essentially the same story.¹⁸

The big problem with using Type I sum of squares is the fact that it really does depend on the order in which you enter the variables. Yet, in many situations the researcher has no reason to prefer one ordering over another. This is presumably the case for our milk and sugar problem. Should we add milk first, or sugar first? It feels exactly as arbitrary as a data analysis question as it does as a coffee-making question. There may in fact be some people with firm opinions about ordering, but it's hard to imagine a principled answer to the question. Yet, look what happens when we change the ordering:

```
> mod <- lm( babble ~ milk + sugar + sugar:milk, coffee )
> anova( mod )
Analysis of Variance Table

Response: babble
  Df Sum Sq Mean Sq F value    Pr(>F)
milk      1 1.4440 1.44400  5.4792 0.037333 *
sugar     2 3.0696 1.53482  5.8238 0.017075 *
milk:sugar 2 5.9439 2.97193 11.2769 0.001754 **
Residuals 12 3.1625 0.26354
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

The *p*-values for both main effect terms have changed, and fairly dramatically. Among other things, the effect of `milk` has become significant (though one should avoid drawing any strong conclusions about this, as I've mentioned previously). Which of these two ANOVAs should one report? It's not immediately obvious.

When you look at the hypothesis tests that are used to define the “first” main effect and the “second” one, it's clear that they're qualitatively different from one another. In our initial example, we saw that the test for the main effect of `sugar` completely ignores `milk`, whereas the test of the main effect of `milk` does take `sugar` into account. As such, the Type I testing strategy really does treat the first main effect as if it had a kind of theoretical primacy over the second one. In my experience there is very rarely if ever any theoretically primacy of this kind that would justify treating any two main effects asymmetrically.

The consequence of all this is that Type I tests are very rarely of much interest, and so we should move on to discuss Type II tests and Type III tests. However, for the sake of completeness – on the off chance that you ever find yourself needing to run Type I tests – I'll comment briefly on how R determines the ordering of terms in a Type I test. The key principle in Type I sum of squares is that the hypothesis testing be sequential, with terms added one at a time. However, it does also imply that main effects be added first (e.g., factors `A`, `B`, `C` etc), followed by first order interaction terms (e.g., terms like `A:B` and `B:C`), then second order interactions (e.g., `A:B:C`) and so on. Within each “block” you can specify whatever order you like. So, for instance, if we specified our model using a command like this,

```
> mod <- lm( outcome ~ A + B + C + B:C + A:B + A:C )
```

and then used `anova(mod)` to produce sequential hypothesis tests, what we'd see is that the main effect terms would be entered `A` then `B` and then `C`, but then the interactions would be entered in the order `B:C` first, then `A:B` and then finally `A:C`. Reordering the terms within each group will change the ordering, as we saw earlier. However, changing the order of terms across blocks has no effect. For instance, if we tried to move the interaction term `B:C` to the front, like this,

¹⁸The one thing that might seem a little opaque to some people is why the residual degrees of freedom in this output look different from one another (i.e., ranging from 12 to 17) whereas in the original one the residual degrees of freedom is fixed at 12. It's actually the case that R uses a residual df of 12 in all cases (that's why the *p* values are the same in the two outputs, and it's enough to verify that `pnorm(6.7495, 2,12, lower.tail=FALSE)` gives the correct answer of *p* = .010863, for instance, whereas `pnorm(6.7495, 2,15, lower.tail=FALSE)` would have given a *p*-value of about .00812. It's the residual degrees of freedom in the full model (i.e., the last one) that matters here.

```
> mod <- lm( outcome ~ B:C + A + B + C + A:B + A:C )
```

it would have no effect. R would still enter the terms in the same order as last time. If for some reason you really, really need an interaction term to be entered first, then you have to do it the long way, creating each model manually using a separate `lm()` command and then using a command like `anova(mod.1, mod.2, mod.3, mod.4)` to force R to enter them in the order that you want.

16.10.4 Type III sum of squares

Having just finished talking about Type I tests, you might think that the natural thing to do next would be to talk about Type II tests. However, I think it's actually a bit more natural to discuss Type III tests (which are simple) before talking about Type II tests (which are trickier). The basic idea behind Type III tests is extremely simple: regardless of which term you're trying to evaluate, run the *F*-test in which the alternative hypothesis corresponds to the full ANOVA model as specified by the user, and the null model just deletes that one term that you're testing. For instance, in the coffee example, in which our full model was `babble ~ sugar + milk + sugar:milk`, the test for a main effect of `sugar` would correspond to a comparison between the following two models:

Null model: `babble ~ milk + sugar:milk`
Alternative model: `babble ~ sugar + milk + sugar:milk`

Similarly the main effect of `milk` is evaluated by testing the full model against a null model that removes the `milk` term, like so:

Null model: `babble ~ sugar + sugar:milk`
Alternative model: `babble ~ sugar + milk + sugar:milk`

Finally, the interaction term `sugar:milk` is evaluated in exactly the same way. Once again, we test the full model against a null model that removes the `sugar:milk` interaction term, like so:

Null model: `babble ~ sugar + milk`
Alternative model: `babble ~ sugar + milk + sugar:milk`

The basic idea generalises to higher order ANOVAs. For instance, suppose that we were trying to run an ANOVA with three factors, **A**, **B** and **C**, and we wanted to consider all possible main effects and all possible interactions, including the three way interaction **A:B:C**. The table below shows you what the Type III tests look like for this situation:

Term being tested is	Null model is <code>outcome ~ ...</code>	Alternative model is <code>outcome ~ ...</code>
A	<code>B + C + A:B + A:C + B:C + A:B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>
B	<code>A + C + A:B + A:C + B:C + A:B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>
C	<code>A + B + A:B + A:C + B:C + A:B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>
A:B	<code>A + B + C + A:C + B:C + A:B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>
A:C	<code>A + B + C + A:B + B:C + A:B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>
B:C	<code>A + B + C + A:B + A:C + A:B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>
A:B:C	<code>A + B + C + A:B + A:C + B:C</code>	<code>A + B + C + A:B + A:C + B:C + A:B:C</code>

As ugly as that table looks, it's pretty simple. In all cases, the alternative hypothesis corresponds to the full model, which contains three main effect terms (e.g. **A**), three first order interactions (e.g. **A:B**) and one second order interaction (i.e., **A:B:C**). The null model always contains 6 of the 7 terms: and the missing one is the one whose significance we're trying to test.

At first pass, Type III tests seem like a nice idea. Firstly, we've removed the asymmetry that caused us to have problems when running Type I tests. And because we're now treating all terms the same way,

the results of the hypothesis tests do not depend on the order in which we specify them. This is definitely a good thing. However, there is a big problem when interpreting the results of the tests, especially for main effect terms. Consider the coffee data. Suppose it turns out that the main effect of `milk` is not significant according to the Type III tests. What this is telling us is that `babble ~ sugar + sugar:milk` is a better model for the data than the full model. But what does that even *mean*? If the interaction term `sugar:milk` was also non-significant, we'd be tempted to conclude that the data are telling us that the only thing that matters is `sugar`. But suppose we have a significant interaction term, but a non-significant main effect of `milk`. In this case, are we to assume that there really is an “effect of sugar”, an “interaction between milk and sugar”, but no “effect of milk”? That seems crazy. The right answer simply *must* be that it's meaningless¹⁹ to talk about the main effect if the interaction is significant. In general, this seems to be what most statisticians advise us to do, and I think that's the right advice. But if it really is meaningless to talk about non-significant main effects in the presence of a significant interaction, then it's not at all obvious why Type III tests should allow the null hypothesis to rely on a model that includes the interaction but omits one of the main effects that make it up. When characterised in this fashion, the null hypotheses really don't make much sense at all.

Later on, we'll see that Type III tests can be redeemed in some contexts, but I'd better show you how to actually compute a Type III ANOVA first. The `anova()` function in R does not directly support Type II tests or Type III tests. Technically, you *can* do it by creating the various models that form the null and alternative hypotheses for each test, and then using `anova()` to compare the models to one another. I outlined the gist of how that would be done when talking about Type I tests, but speaking from first hand experience²⁰ I can tell you that it's very tedious. In practice, the `anova()` function is only used to produce Type I tests or to compare specific models of particular interest (see Section 16.5). If you want Type II or Type III tests you need to use the `Anova()` function in the `car` package. It's pretty easy to use, since there's a `type` argument that you specify. So, to return to our coffee example, our Type III tests are run as follows:

```
> mod <- lm( babble ~ sugar * milk, coffee )
> Anova( mod, type=3 )
Anova Table (Type III tests)

Response: babble
      Sum Sq Df F value    Pr(>F)
(Intercept) 41.070  1 155.839 3.11e-08 ***
sugar        5.880  2 11.156 0.001830 **
milk         4.107  1 15.584 0.001936 **
sugar:milk   5.944  2 11.277 0.001754 **
Residuals    3.162 12
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

As you can see, I got lazy this time and used `sugar * milk` as a shorthand way of referring to `sugar + milk + sugar:milk`. The important point here is that this is just a regular ANOVA table, and we can see that our Type III tests are significant for all terms, even the intercept.

Except, as usual, it's not that simple. One of the perverse features of the Type III testing strategy is that the results turn out to depend on the *contrasts* that you use to encode your factors (see Section 16.7 if you've forgotten what the different types of contrasts are). The results that I presented in the ANOVA

¹⁹Or, at the very least, rarely of interest.

²⁰Yes, I'm actually a big enough nerd that I've written my own functions implementing Type II tests and Type III tests. I only did it to convince myself that I knew how the different Types of test worked, but it did turn out to be a handy exercise: the `etaSquared()` function in the `lsr` package relies on it. There's actually even an argument in the `etaSquared()` function called `anova`. By default, `anova=FALSE` and the function just prints out the effect sizes. However, if you set `anova=TRUE` it will spit out the full ANOVA table as well. This works for Types I, II and III. Just set the `types` argument to select which type of test you want.

table above are based on the R default, which is treatment contrasts; and as we'll see later, this is usually a very poor choice if you want to run Type III tests. So let's see what happens if switch to Helmert contrasts:

```
> my.contrasts <- list( milk = "contr.Helmert", sugar = "contr.Helmert" )
> mod.H <- lm( babble ~ sugar * milk, coffee, contrasts = my.contrasts )
> Anova( mod.H, type=3 )
Anova Table (Type III tests)

Response: babble
      Sum Sq Df  F value    Pr(>F)
(Intercept) 434.29  1 1647.8882 3.231e-14 ***
sugar        2.13   2    4.0446  0.045426 *
milk         1.00   1    3.8102  0.074672 .
sugar:milk   5.94   2   11.2769  0.001754 **
Residuals    3.16  12
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

Oh, that's not good at all. In the case of `milk` in particular, the *p*-value has changed from .002 to .07. This is a pretty substantial difference, and hopefully it gives you a sense of how important it is that you take care when using Type III tests.

Okay, so if the *p*-values that come out of Type III analyses are so sensitive to the choice of contrasts, does that mean that Type III tests are essentially arbitrary and not to be trusted? To some extent that's true, and when we turn to a discussion of Type II tests we'll see that Type II analyses avoid this arbitrariness entirely, but I think that's too strong a conclusion. Firstly, it's important to recognise that some choices of contrasts will always produce the same answers. Of particular importance is the fact that if the columns of our contrast matrix are all constrained to sum to zero, then the Type III analysis will always give the same answers. This means that you'll get the same answers if you use `contr.Helmert` or `contr.sum` or `contr.poly`, but different answers for `contr.treatment` or `contr.SAS`.

```
> random.contrasts <- matrix( rnorm(6), 3, 2 )  # create a random matrix
> random.contrasts[, 1] <- random.contrasts[, 1] - mean( random.contrasts[, 1] ) # contrast 1 sums to 0
> random.contrasts[, 2] <- random.contrasts[, 2] - mean( random.contrasts[, 2] ) # contrast 2 sums to 0
> random.contrasts  # print it to check that we really have an arbitrary contrast matrix...
     [,1]      [,2]
[1,]  0.38898807 -0.78454935
[2,] -0.04337123  0.70004953
[3,] -0.34561683  0.08449982

> contrasts( coffee$sugar ) <- random.contrasts # random contrasts for sugar
> contrasts( coffee$milk ) <- contr.Helmert(2) # Helmert contrasts for the milk factor
> mod.R <- lm( babble ~ sugar * milk, coffee ) # R will use the contrasts that we assigned
> Anova( mod.R, type = 3 )
Anova Table (Type III tests)

Response: babble
      Sum Sq Df  F value    Pr(>F)
(Intercept) 434.29  1 1647.8882 3.231e-14 ***
sugar        2.13   2    4.0446  0.045426 *
milk         1.00   1    3.8102  0.074672 .
sugar:milk   5.94   2   11.2769  0.001754 **
Residuals    3.16  12
```

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

Yep, same answers.

16.10.5 Type II sum of squares

Okay, so we've seen Type I and III tests now, and both are pretty straightforward: Type I tests are performed by gradually adding terms one at a time, whereas Type III tests are performed by taking the full model and looking to see what happens when you remove each term. However, both have some serious flaws: Type I tests are dependent on the order in which you enter the terms, and Type III tests are dependent on how you code up your contrasts. Because of these flaws, neither one is easy to interpret. Type II tests are a little harder to describe, but they avoid both of these problems, and as a result they are a little easier to interpret.

Type II tests are broadly similar to Type III tests: start with a "full" model, and test a particular term by deleting it from that model. However, Type II tests are based on the **marginality principle** which states that you should not omit a lower order term from your model if there are any higher order ones that depend on it. So, for instance, if your model contains the interaction **A:B** (a 2nd order term), then it really ought to contain the main effects **A** and **B** (1st order terms). Similarly, if it contains a three way interaction term **A:B:C**, then the model must also include the main effects **A**, **B** and **C** as well as the simpler interactions **A:B**, **A:C** and **B:C**. Type III tests routinely violate the marginality principle. For instance, consider the test of the main effect of **A** in the context of a three-way ANOVA that includes all possible interaction terms. According to Type III tests, our null and alternative models are:

Null model: **outcome ~ B + C + A:B + A:C + B:C + A:B:C**
Alternative model: **outcome ~ A + B + C + A:B + A:C + B:C + A:B:C**

Notice that the null hypothesis omits **A**, but includes **A:B**, **A:C** and **A:B:C** as part of the model. This, according to the Type II tests, is not a good choice of null hypothesis. What we should do instead, if we want to test the null hypothesis that **A** is not relevant to our **outcome**, is to specify the null hypothesis that is the most complicated model that does not rely on **A** in any form, even as an interaction. The alternative hypothesis corresponds to this null model plus a main effect term of **A**. This is a lot closer to what most people would intuitively think of as a "main effect of **A**", and it yields the following as our Type II test of the main effect of **A**.²¹

Null model: **outcome ~ B + C + B:C**
Alternative model: **outcome ~ A + B + C + B:C**

Anyway, just to give you a sense of how the Type II tests play out, here's the full table of tests that would be applied in a three-way factorial ANOVA:

²¹Note, of course, that this does depend on the model that the user specified. If original ANOVA model doesn't contain an interaction term for **B:C**, then obviously it won't appear in either the null or the alternative. But that's true for Types I, II and III. They never include any terms that you *didn't* include, but they make different choices about how to construct tests for the ones that you did include.

Term being tested is	Null model is <code>outcome ~ ...</code>	Alternative model is <code>outcome ~ ...</code>
A	B + C + B:C	A + B + C + B:C
B	A + C + A:C	A + B + C + A:C
C	A + B + A:B	A + B + C + A:B
A:B	A + B + C + A:C + B:C	A + B + C + A:B + A:C + B:C
A:C	A + B + C + A:B + B:C	A + B + C + A:B + A:C + B:C
B:C	A + B + C + A:B + A:C	A + B + C + A:B + A:C + B:C
A:B:C	A + B + C + A:B + A:C + B:C	A + B + C + A:B + A:C + B:C + A:B:C

In the context of the two way ANOVA that we've been using in the coffee data, the hypothesis tests are even simpler. The main effect of `sugar` corresponds to an *F*-test comparing these two models:

$$\begin{aligned} \text{Null model: } & \text{babble} \sim \text{milk} \\ \text{Alternative model: } & \text{babble} \sim \text{sugar} + \text{milk} \end{aligned}$$

The test for the main effect of `milk` is

$$\begin{aligned} \text{Null model: } & \text{babble} \sim \text{sugar} \\ \text{Alternative model: } & \text{babble} \sim \text{sugar} + \text{milk} \end{aligned}$$

Finally, the test for the interaction `sugar:milk` is:

$$\begin{aligned} \text{Null model: } & \text{babble} \sim \text{sugar} + \text{milk} \\ \text{Alternative model: } & \text{babble} \sim \text{sugar} + \text{milk} + \text{sugar:milk} \end{aligned}$$

Running the tests are again straightforward. We use the `Anova()` function, specifying `type=2`:

```
> mod <- lm( babble ~ sugar*milk, coffee )
> Anova( mod, type = 2 )
Anova Table (Type II tests)

Response: babble
           Sum Sq Df F value    Pr(>F)
sugar      3.0696  2  5.8238 0.017075 *
milk       0.9561  1  3.6279 0.081061 .
sugar:milk 5.9439  2 11.2769 0.001754 **
Residuals  3.1625 12
---
Signif. codes:  0 *** 0.001 ** 0.01 * 0.05 . 0.1   1
```

Type II tests have some clear advantages over Type I and Type III tests. They don't depend on the order in which you specify factors (unlike Type I), and they don't depend on the contrasts that you use to specify your factors (unlike Type III). And although opinions may differ on this last point, and it will definitely depend on what you're trying to do with your data, I do think that the hypothesis tests that they specify are more likely to correspond to something that you actually care about. As a consequence, I find that it's usually easier to interpret the results of a Type II test than the results of a Type I or Type III test. For this reason, my tentative advice is that, if you can't think of any obvious model comparisons that directly map onto your research questions but you still want to run an ANOVA in an unbalanced

design, Type II tests are probably a better choice than Type I or Type III.²²

16.10.6 Effect sizes (and non-additive sums of squares)

The `etaSquared()` function in the `lsr` package computes η^2 and partial η^2 values for unbalanced designs and for different Types of tests. It's pretty straightforward. All you have to do is indicate which `type` of tests you're doing,

```
> etaSquared( mod, type=2 )
      eta.sq eta.sq.part
sugar     0.22537682  0.4925493
milk      0.07019886  0.2321436
sugar:milk 0.43640732  0.6527155
```

and out pops the η^2 and partial η^2 values, as requested. However, when you've got an unbalanced design, there's a bit of extra complexity involved. To see why, let's expand the output from the `etaSquared()` function so that it displays the full ANOVA table:

```
> es <- etaSquared( mod, type=2, anova=TRUE )
> es
      eta.sq eta.sq.part      SS df      MS      F      p
sugar     0.22537682  0.4925493 3.0696323  2 1.5348161  5.823808 0.017075099
milk      0.07019886  0.2321436 0.9561085  1 0.9561085  3.627921 0.081060698
sugar:milk 0.43640732  0.6527155 5.9438677  2 2.9719339 11.276903 0.001754333
Residuals 0.23219530          NA 3.1625000 12 0.2635417          NA          NA
```

Okay, if you remember back to our very early discussions of ANOVA, one of the key ideas behind the sums of squares calculations is that if we add up all the SS terms associated with the effects in the model, and add that to the residual SS, they're supposed to add up to the total sum of squares. And, on top of that, the whole idea behind η^2 is that – because you're dividing one of the SS terms by the total SS value – is that an η^2 value can be interpreted as the proportion of variance accounted for by a particular term.

Now take a look at the output above. Because I've included the η^2 value associated with the residuals (i.e., proportion of variance in the outcome attributed to the residuals, rather than to one of the effects), you'd expect all the η^2 values to sum to 1. Because, the whole idea here was that the variance in the outcome variable can be divided up into the variability attributable to the model, and the variability in the residuals. Right? Right? And yet when we add up the η^2 values for our model...

```
> sum( es[, "eta.sq"] )
[1] 0.9641783
```

... we discover that for Type II and Type III tests they generally don't sum to 1. Some of the variability has gone “missing”. It's not being attributed to the model, and it's not being attributed to the residuals either. What's going on here?

²²I find it amusing to note that the default in R is Type I and the default in SPSS is Type III (with Helmert contrasts). Neither of these appeals to me all that much. Relatedly, I find it depressing that almost nobody in the psychological literature ever bothers to report which Type of tests they ran, much less the order of variables (for Type I) or the contrasts used (for Type III). Often they don't report what software they used either. The only way I can ever make any sense of what people typically report is to try to guess from auxiliary cues which software they were using, and to assume that they never changed the default settings. Please don't do this... now that you know about these issues, make sure you indicate what software you used, and if you're reporting ANOVA results for unbalanced data, then specify what Type of tests you ran, specify order information if you've done Type I tests and specify contrasts if you've done Type III tests. Or, even better, do hypotheses tests that correspond to things you really care about, and then report those!

Before giving you the answer, I want to push this idea a little further. From a mathematical perspective, it's easy enough to see that the missing variance is a consequence of the fact that in Types II and III, the individual SS values are not obliged to the total sum of squares, and will only do so if you have balanced data. I'll explain why this happens and what it means in a second, but first let's verify that this is the case using the ANOVA table. First, we can calculate the total sum of squares directly from the raw data:

```
> ss.tot <- sum( (coffee$babble - mean(coffee$babble))^2 )
> ss.tot
[1] 13.62
```

Next, we can read off all the SS values from one of our Type I ANOVA tables, and add them up. As you can see, this gives us the same answer, just like it's supposed to:

```
> type.I.sum <- 3.5575 + 0.9561 + 5.9439 + 3.1625
> type.I.sum
[1] 13.62
```

However, when we do the same thing for the Type II ANOVA table, it turns out that the SS values in the table add up to slightly less than the total SS value:

```
> type.II.sum <- 0.9561 + 3.0696 + 5.9439 + 3.1625
> type.II.sum
[1] 13.1321
```

So, once again, we can see that there's a little bit of variance that has "disappeared" somewhere.

Okay, time to explain what's happened. The reason why this happens is that, when you have unbalanced designs, your factors become correlated with one another, and it becomes difficult to tell the difference between the effect of Factor A and the effect of Factor B. In the extreme case, suppose that we'd run a 2×2 design in which the number of participants in each group had been as follows:

	sugar	no sugar
milk	100	0
no milk	0	100

Here we have a spectacularly unbalanced design: 100 people have milk and sugar, 100 people have no milk and no sugar, and that's all. There are 0 people with milk and no sugar, and 0 people with sugar but no milk. Now suppose that, when we collected the data, it turned out there is a large (and statistically significant) difference between the "milk and sugar" group and the "no-milk and no-sugar" group. Is this a main effect of sugar? A main effect of milk? Or an interaction? It's impossible to tell, because the presence of sugar has a perfect association with the presence of milk. Now suppose the design had been a little more balanced:

	sugar	no sugar
milk	100	5
no milk	5	100

This time around, it's technically possible to distinguish between the effect of milk and the effect of sugar, because we have a few people that have one but not the other. However, it will still be pretty difficult to do so, because the association between sugar and milk is still extremely strong, and there are so few observations in two of the groups. Again, we're very likely to be in the situation where we *know* that the predictor variables (milk and sugar) are related to the outcome (babbling), but we don't know if the *nature* of that relationship is a main effect of one predictor, or the other predictor or the interaction.

This uncertainty is the reason for the missing variance. The “missing” variance corresponds to variation in the outcome variable that is clearly attributable to the predictors, but we don’t know which of the effects in the model is responsible. When you calculate Type I sum of squares, no variance ever goes missing: the sequential nature of Type I sum of squares means that the ANOVA automatically attributes this variance to whichever effects are entered first. However, the Type II and Type III tests are more conservative. Variance that cannot be clearly attributed to a specific effect doesn’t get attributed to any of them, and it goes missing.

16.11

Summary

- Factorial ANOVA with balanced designs, without interactions (Section 16.1) and with interactions included (Section 16.2)
- Effect size, estimated means, and confidence intervals in a factorial ANOVA (Section 16.3)
- Understanding the linear model underlying ANOVA (Sections 16.5, 16.6 and 16.7)
- Post hoc testing using Tukey’s HSD (Section 16.8), and a brief commentary on planned comparisons (Section 16.9)
- Factorial ANOVA with unbalanced designs (Section 16.10)

17. Epilogue

It feels somewhat strange to be writing this chapter, and more than a little inappropriate. An epilogue is what you write when a book is finished, and this book really isn't finished. There are a lot of things still missing from this book. It doesn't have an index yet. The chapter "summaries" are usually no more than a list of section titles. A *lot* of references are missing. There are no "do it yourself" exercises. And in general, I feel that there a lot of things that are wrong with the presentation, organisation and content of this book. Given all that, I don't want to try to write a "proper" epilogue. I haven't finished writing the substantive content yet, so it doesn't make sense to try to bring it all together. But this version of the book is going to go online for students to use, and you will be able to purchase a hard copy too, so I want to give it at least a veneer of closure. So let's give it a go, shall we?

17.1

The undiscovered statistics

First, I'm going to talk a bit about some of the content that I wish I'd had the chance to cram into this version of the book, just so that you can get a sense of what other ideas are out there in the world of statistics. I think this would be important even if this book were getting close to a final product: one thing that students often fail to realise is that their introductory statistics classes are just that: an introduction. If you want to go out into the wider world and do real data analysis, you have to learn a whole lot of new tools that extend the content of your undergraduate lectures in all sorts of different ways. Don't assume that something can't be done just because it wasn't covered in undergrad. Don't assume that something is the right thing to do just because it *was* covered in an undergrad class. To stop you from falling victim to that trap, I think it's useful to give a bit of an overview of some of the other ideas out there.

17.1.1 Omissions within the topics covered

Even within the topics that I have covered in the book, there are a lot of omissions that I'd like to redress in future version of the book. Just sticking to things that are purely about statistics (rather than things associated with R), the following is a representative but not exhaustive list of topics that I'd like to expand on in later versions:

- **Other types of correlations.** In Chapter 5 I talked about two types of correlation: Pearson and Spearman. Both of these methods of assessing correlation are applicable to the case where you have two continuous variables and want to assess the relationship between them. What about the case where your variables are both nominal scale? Or when one is nominal scale and the other is

continuous? There are actually methods for computing correlations in such cases (e.g., polychoric correlation), but I just haven't had time to write about them yet.

- **More detail on effect sizes.** In general, I think the treatment of effect sizes throughout the book is a little more cursory than it should be. In almost every instance, I've tended just to pick one measure of effect size (usually the most popular one) and describe that. However, for almost all tests and models there are multiple ways of thinking about effect size, and I'd like to go into more detail in the future.
- **Dealing with violated assumptions.** In a number of places in the book I've talked about some things you can do when you find that the assumptions of your test (or model) are violated, but I think that I ought to say more about this. In particular, I think it would have been nice to talk in a lot more detail about how you can transform variables to fix problems. I talked a bit about this in Sections 7.2, 7.3 and 15.8.4, but the discussion isn't detailed enough I think.
- **Interaction terms for regression.** In Chapter 16 I talked about the fact that you can have interaction terms in an ANOVA, and I also pointed out that ANOVA can be interpreted as a kind of linear regression model. Yet, when talking about regression in Chapter 15 I made not mention of interactions at all. However, there's nothing stopping you from including interaction terms in a regression model. It's just a little more complicated to figure out what an "interaction" actually means when you're talking about the interaction between two continuous predictors, and it can be done in more than one way. Even so, I would have liked to talk a little about this.
- **Method of planned comparison.** As I mentioned this in Chapter 16, it's not always appropriate to be using post hoc correction like Tukey's HSD when doing an ANOVA, especially when you had a very clear (and limited) set of comparisons that you cared about ahead of time. I would like to talk more about this in a future version of book.
- **Multiple comparison methods.** Even within the context of talking about post hoc tests and multiple comparisons, I would have liked to talk about the methods in more detail, and talk about what other methods exist besides the few options I mentioned.

17.1.2 Statistical models missing from the book

Statistics is a huge field. The core tools that I've described in this book (chi-square tests, *t*-tests, ANOVA and regression) are basic tools that are widely used in everyday data analysis, and they form the core of most introductory stats books. However, there are a lot of other tools out there. There are so very many data analysis situations that these tools don't cover, and in future versions of this book I want to talk about them. To give you a sense of just how much more there is, and how much more work I want to do to finish this thing, the following is a list of statistical modelling tools that I would have liked to talk about. Some of these will definitely make it into future versions of the book.

- **Analysis of covariance.** In Chapter 16 I spent a bit of time discussing the connection between ANOVA and regression, pointing out that any ANOVA model can be recast as a kind of regression model. More generally, both are examples of *linear models*, and it's quite possible to consider linear models that are more general than either. The classic example of this is "analysis of covariance" (ANCOVA), and it refers to the situation where some of your predictors are continuous (like in a regression model) and others are categorical (like in an ANOVA).
- **Nonlinear regression.** When discussing regression in Chapter 15, we saw that regression assume that the relationship between predictors and outcomes is linear. On the other hand, when we talked about the simpler problem of correlation in Chapter 5, we saw that there exist tools (e.g.,

Spearman correlations) that are able to assess non-linear relationships between variables. There are a number of tools in statistics that can be used to do non-linear regression. For instance, some non-linear regression models assume that the relationship between predictors and outcomes is monotonic (e.g., isotonic regression), while others assume that it is smooth but not necessarily monotonic (e.g., Lowess regression), while others assume that the relationship is of a known form that happens to be nonlinear (e.g., polynomial regression).

- **Logistic regression.** Yet another variation on regression occurs when the outcome variable is binary valued, but the predictors are continuous. For instance, suppose you’re investigating social media, and you want to know if it’s possible to predict whether or not someone is on Twitter as a function of their income, their age, and a range of other variables. This is basically a regression model, but you can’t use regular linear regression because the outcome variable is binary (you’re either on Twitter or you’re not): because the outcome variable is binary, there’s no way that the residuals could possibly be normally distributed. There are a number of tools that statisticians can apply to this situation, the most prominent of which is logistic regression.
- **The General Linear Model (GLM).** The GLM is actually a family of models that includes logistic regression, linear regression, (some) nonlinear regression, ANOVA and many others. The basic idea in the GLM is essentially the same idea that underpins linear models, but it allows for the idea that your data might not be normally distributed, and allows for nonlinear relationships between predictors and outcomes. There are a lot of very handy analyses that you can run that fall within the GLM, so it’s a very useful thing to know about.
- **Survival analysis.** In Chapter 2 I talked about “differential attrition”, the tendency for people to leave the study in a non-random fashion. Back then, I was talking about it as a potential methodological concern, but there are a lot of situations in which differential attrition is actually the thing you’re interested in. Suppose, for instance, you’re interested in finding out how long people play different kinds of computer games in a single session. Do people tend to play RTS (real time strategy) games for longer stretches than FPS (first person shooter) games? You might design your study like this. People come into the lab, and they can play for as long or as little as they like. Once they’re finished, you record the time they spent playing. However, due to ethical restrictions, let’s suppose that you cannot allow them to keep playing longer than two hours. A lot of people will stop playing before the two hour limit, so you know exactly how long they played. But some people will run into the two hour limit, and so you don’t know how long they would have kept playing if you’d been able to continue the study. As a consequence, your data are systematically *censored*: you’re missing all of the very long times. How do you analyse this data sensibly? This is the problem that survival analysis solves. It is specifically designed to handle this situation, where you’re systematically missing one “side” of the data because the study ended. It’s very widely used in health research, and in that context it is often literally used to analyse survival. For instance, you may be tracking people with a particular type of cancer, some who have received treatment A and others who have received treatment B, but you only have funding to track them for 5 years. At the end of the study period some people are alive, others are not. In this context, survival analysis is useful for determining which treatment is more effective, and telling you about the risk of death that people face over time.
- **Repeated measures ANOVA.** When talking about reshaping data in Chapter 7, I introduced some data sets in which each participant was measured in multiple conditions (e.g., in the drugs data set, the working memory capacity (WMC) of each person was measured under the influence of alcohol and caffeine). It is quite common to design studies that have this kind of repeated measures structure. A regular ANOVA doesn’t make sense for these studies, because the repeated measurements mean that independence is violated (i.e., observations from the same participant are more closely related to one another than to observations from other participants). Repeated measures ANOVA is a tool that can be applied to data that have this structure. The basic idea

behind RM-ANOVA is to take into account the fact that participants can have different overall levels of performance. For instance, Amy might have a WMC of 7 normally, which falls to 5 under the influence of caffeine, whereas Borat might have a WMC of 6 normally, which falls to 4 under the influence of caffeine. Because this is a repeated measures design, we recognise that – although Amy has a higher WMC than Borat – the effect of caffeine is identical for these two people. In other words, a repeated measures design means that we can attribute some of the variation in our WMC measurement to individual differences (i.e., some of it is just that Amy has higher WMC than Borat), which allows us to draw stronger conclusions about the effect of caffeine.

- **Mixed models.** Repeated measures ANOVA is used in situations where you have observations clustered within experimental units. In the example I gave above, we have multiple WMC measures for each participant (i.e., one for each condition). However, there are a lot of other ways in which you can end up with multiple observations per participant, and for most of those situations the repeated measures ANOVA framework is insufficient. A good example of this is when you track individual people across multiple time points. Let's say you're tracking happiness over time, for two people. Aaron's happiness starts at 10, then drops to 8, and then to 6. Belinda's happiness starts at 6, then rises to 8 and then to 10. Both of these two people have the same "overall" level of happiness (the average across the three time points is 8), so a repeated measures ANOVA analysis would treat Aaron and Belinda the same way. But that's clearly wrong. Aaron's happiness is decreasing, whereas Belinda's is increasing. If you want to optimally analyse data from an experiment where people can change over time, then you need a more powerful tool than repeated measures ANOVA. The tools that people use to solve this problem are called "mixed" models, because they are designed to learn about individual experimental units (e.g. happiness of individual people over time) as well as overall effects (e.g. the effect of money on happiness over time). Repeated measures ANOVA is perhaps the simplest example of a mixed model, but there's a lot you can do with mixed models that you can't do with repeated measures ANOVA.
- **Reliability analysis.** Back in Chapter 2 I talked about reliability as one of the desirable characteristics of a measurement. One of the different types of reliability I mentioned was inter-item reliability. For example, when designing a survey used to measure some aspect to someone's personality (e.g., extraversion), one generally attempts to include several different questions that all ask the same basic question in lots of different ways. When you do this, you tend to expect that all of these questions will tend to be correlated with one another, because they're all measuring the same latent construct. There are a number of tools (e.g., Cronbach's α) that you can use to check whether this is actually true for your study.
- **Factor analysis.** One big shortcoming with reliability measures like Cronbach's α is that they assume that your observed variables are all measuring a *single* latent construct. But that's not true in general. If you look at most personality questionnaires, or IQ tests, or almost anything where you're taking lots of measurements, it's probably the case that you're actually measuring several things at once. For example, all the different tests used when measuring IQ do tend to correlate with one another, but the pattern of correlations that you see across tests suggests that there are multiple different "things" going on in the data. Factor analysis (and related tools like principal components analysis and independent components analysis) is a tool that you can use to help you figure out what these things are. Broadly speaking, what you do with these tools is take a big correlation matrix that describes all pairwise correlations between your variables, and attempt to express this pattern of correlations using only a small number of latent variables. Factor analysis is a very useful tool – it's a great way of trying to see how your variables are related to one another – but it can be tricky to use well. A lot of people make the mistake of thinking that when factor analysis uncovers a latent variable (e.g., extraversion pops out as a latent variable when you factor analyse most personality questionnaires), it must actually correspond to a real "thing". That's not necessarily true. Even so, factor analysis is a very useful thing to know about (especially for psychologists), and I do want to talk about it in a later version of the book.

- **Multidimensional scaling.** Factor analysis is an example of an “unsupervised learning” model. What this means is that, unlike most of the “supervised learning” tools I’ve mentioned, you can’t divide up your variables into predictors and outcomes. Regression is supervised learning; factor analysis is unsupervised learning. It’s not the only type of unsupervised learning model however. For example, in factor analysis one is concerned with the analysis of correlations between variables. However, there are many situations where you’re actually interested in analysing similarities or dissimilarities between objects, items or people. There are a number of tools that you can use in this situation, the best known of which is multidimensional scaling (MDS). In MDS, the idea is to find a “geometric” representation of your items. Each item is “plotted” as a point in some space, and the distance between two points is a measure of how dissimilar those items are.
- **Clustering.** Another example of an unsupervised learning model is clustering (also referred to as classification), in which you want to organise all of your items into meaningful groups, such that similar items are assigned to the same groups. A lot of clustering is unsupervised, meaning that you don’t know anything about what the groups are, you just have to guess. There are other “supervised clustering” situations where you need to predict group memberships on the basis of other variables, and those group memberships are actually observables: logistic regression is a good example of a tool that works this way. However, when you don’t actually know the group memberships, you have to use different tools (e.g., k -means clustering). There’s even situations where you want to do something called “semi-supervised clustering”, in which you know the group memberships for some items but not others. As you can probably guess, clustering is a pretty big topic, and a pretty useful thing to know about.
- **Causal models.** One thing that I haven’t talked about much in this book is how you can use statistical modeling to learn about the causal relationships between variables. For instance, consider the following three variables which might be of interest when thinking about how someone died in a firing squad. We might want to measure whether or not an execution order was given (variable A), whether or not a marksman fired their gun (variable B), and whether or not the person got hit with a bullet (variable C). These three variables are all correlated with one another (e.g., there is a correlation between guns being fired and people getting hit with bullets), but we actually want to make stronger statements about them than merely talking about correlations. We want to talk about causation. We want to be able to say that the execution order (A) causes the marksman to fire (B) which causes someone to get shot (C). We can express this by a directed arrow notation: we write it as $A \rightarrow B \rightarrow C$. This “causal chain” is a fundamentally different explanation for events than one in which the marksman fires first, which causes the shooting $B \rightarrow C$, and then causes the executioner to “retroactively” issue the execution order, $B \rightarrow A$. This “common effect” model says that A and C are both caused by B. You can see why these are different. In the first causal model, if we had managed to stop the executioner from issuing the order (intervening to change A), then no shooting would have happened. In the second model, the shooting would have happened any way because the marksman was *not* following the execution order. There is a big literature in statistics on trying to understand the causal relationships between variables, and a number of different tools exist to help you test different causal stories about your data. The most widely used of these tools (in psychology at least) is structural equations modelling (SEM), and at some point I’d like to extend the book to talk about it.

Of course, even this listing is incomplete. I haven’t mentioned time series analysis, item response theory, market basket analysis, classification and regression trees, or any of a huge range of other topics. However, the list that I’ve given above is essentially my wish list for this book. Sure, it would double the length of the book, but it would mean that the scope has become broad enough to cover most things that applied researchers in psychology would need to use.

17.1.3 Other ways of doing inference

A different sense in which this book is incomplete is that it focuses pretty exclusively on a very narrow and old-fashioned view of how inferential statistics should be done. In Chapter 10 I talked a little bit about the idea of unbiased estimators, sampling distributions and so on. In Chapter 11 I talked about the theory of null hypothesis significance testing and p -values. These ideas have been around since the early 20th century, and the tools that I've talked about in the book rely very heavily on the theoretical ideas from that time. I've felt obligated to stick to those topics because the vast majority of data analysis in science is also reliant on those ideas. However, the theory of statistics is not restricted to those topics, and – while everyone should know about them because of their practical importance – in many respects those ideas do not represent best practice for contemporary data analysis. Here are some of the ideas that I've omitted, all of which I intend to discuss in more detail in future versions of the book:

- **Bayesian methods.** In a few places in the book I've mentioned the fact that there is this thing called “Bayesian probability”, in which probability is interpreted as a “degree of belief” in some proposition (e.g., the proposition that I will get a six when I roll a die). Unlike the conventional frequentist definition of probability, the Bayesian interpretation of probability allows you to assign probability to “one off” events, rather than restricting probability to events that can be replicated. The difference between frequentist and Bayesian interpretations of probability is not purely a matter of interpretation. Bayesian probability leads to different tools for analysing data: if you believe in Bayesian probability, you get different answers for a lot of statistical questions, and (in my opinion) the answers you get are generally better ones. I've talked a little about Bayesian ideas in this book, but Bayesian inference is a huge part of contemporary statistical practice, much moreso than you'd think given the very cursory treatment I've given it here.
- **Bootstrapping.** Throughout the book, whenever I've been introduced a hypothesis test, I've had a strong tendency just to make assertions like “the sampling distribution for BLAH is a t -distribution” or something like that. In some cases, I've actually attempted to justify this assertion. For example, when talking about χ^2 tests in Chapter 12, I made reference to the known relationship between normal distributions and χ^2 distributions (see Chapter 9) to explain how we end up assuming that the sampling distribution of the goodness of fit statistic is χ^2 . However, it's also the case that a lot of these sampling distributions are, well, wrong. The χ^2 test is a good example: it is based on an assumption about the distribution of your data, an assumption which is known to be wrong for small sample sizes! Back in the early 20th century, there wasn't much you could do about this situation: statisticians had developed mathematical results that said that “under assumptions BLAH about the data, the sampling distribution is approximately BLAH”, and that was about the best you could do. A lot of times they didn't even have that: there are lots of data analysis situations for which no-one has found a mathematical solution for the sampling distributions that you need. And so up until the late 20th century, the corresponding tests didn't exist or didn't work. However, computers have changed all that now. There are lots of fancy tricks, and some not-so-fancy, that you can use to get around it. The simplest of these is bootstrapping, and in its simplest form it's incredibly simple. Here it is: simulate the results of your experiment lots and lots of time, under the twin assumptions that (a) the null hypothesis is true and (b) the unknown population distribution actually looks pretty similar to your raw data. In other words, instead of assuming that the data are (for instance) normally distributed, just assume that the population looks the same as your sample, and then use computers to simulate the sampling distribution for your test statistic if that assumption holds. Despite relying on a somewhat dubious assumption (i.e., the population distribution is the same as the sample!) bootstrapping is quick and easy method that works remarkably well in practice for lots of data analysis problems.
- **Cross validation.** One question that pops up in my stats classes every now and then, usually by a student trying to be provocative, is “Why do we care about inferential statistics at all? Why not

just describe your sample?” The answer to the question is usually something like this: “Because our true interest as scientists is not the specific sample that we have observed in the past, we want to make predictions about data we might observe in the future”. A lot of the issues in statistical inference arise because of the fact that we always expect the future to be similar to but a bit different from the past. Or, more generally, new data won’t be quite the same as old data. What we do, in a lot of situations, is try to derive mathematical rules that help us to draw the inferences that are most likely to be correct for new data, rather than to pick the statements that best describe old data. For instance, given two models A and B, and a data set X you collected today, try to pick the model that will best describe a new data set Y that you’re going to collect tomorrow. Sometimes it’s convenient to simulate the process, and that’s what cross-validation does. What you do is divide your data set into two subsets, X1 and X2. Use the subset X1 to train the model (e.g., estimate regression coefficients, let’s say), but then assess the model performance on the other one X2. This gives you a measure of how well the model *generalises* from an old data set to a new one, and is often a better measure of how good your model is than if you just fit it to the full data set X.

- **Robust statistics.** Life is messy, and nothing really works the way it’s supposed to. This is just as true for statistics as it is for anything else, and when trying to analyse data we’re often stuck with all sorts of problems in which the data are just messier than they’re supposed to be. Variables that are supposed to be normally distributed are not *actually* normally distributed, relationships that are supposed to be linear are not *actually* linear, and some of the observations in your data set are almost certainly junk (i.e., not measuring what they’re supposed to). All of this messiness is ignored in most of the statistical theory I developed in this book. However, ignoring a problem doesn’t always solve it. Sometimes, it’s actually okay to ignore the mess, because some types of statistical tools are “robust”: if the data don’t satisfy your theoretical assumptions, they still work pretty well. Other types of statistical tools are not robust: even minor deviations from the theoretical assumptions cause them to break. Robust statistics is a branch of stats concerned with this question, and they talk about things like the “breakdown point” of a statistic: that is, how messy does your data have to be before the statistic cannot be trusted? I touched on this in places. The mean is *not* a robust estimator of the central tendency of a variable; the median is. For instance, suppose I told you that the ages of my five best friends are 34, 39, 31, 43 and 4003 years. How old do you think they are on average? That is, what is the true population mean here? If you use the sample mean as your estimator of the population mean, you get an answer of 830 years. If you use the sample median as the estimator of the population mean, you get an answer of 39 years. Notice that, even though you’re “technically” doing the wrong thing in the second case (using the median to estimate the mean!) you’re actually getting a better answer. The problem here is that one of the observations is clearly, obviously a lie. I don’t have a friend aged 4003 years. It’s probably a typo: I probably meant to type 43. But what if I had typed 53 instead of 43, or 34 instead of 43? Could you be sure if this was a typo? Sometimes the errors in the data are subtle, so you can’t detect them just by eyeballing the sample, but they’re still errors that contaminate your data, and they still affect your conclusions. Robust statistics is a concerned with how you can make *safe* inferences even when faced with contamination that you don’t know about. It’s pretty cool stuff.

17.1.4 Miscellaneous topics

- **Missing data.** Suppose you’re doing a survey, and you’re interested in exercise and weight. You send data to four people. Adam says he exercises a lot and is not overweight. Briony says she exercises a lot and is not overweight. Carol says she does not exercise and is overweight. Dan says he does not exercise and refuses to answer the question about his weight. Elaine does not return the survey. You now have a missing data problem. There is one entire survey missing, and one question missing from another one, What do you do about it? I’ve only barely touched on

this question in this book, in Section 5.8, and in that section all I did was tell you about some R commands you can use to ignore the missing data. But ignoring missing data is not, in general, a safe thing to do. Let's think about Dan's survey here. Firstly, notice that, on the basis of my other responses, I appear to be more similar to Carol (neither of us exercise) than to Adam or Briony. So if you were forced to guess my weight, you'd guess that I'm closer to her than to them. Maybe you'd make some correction for the fact that Adam and I are males and Briony and Carol are females. The statistical name for this kind of guessing is "imputation". Doing imputation safely is hard, but important, especially when the missing data are missing in a systematic way. Because of the fact that people who are overweight are often pressured to feel poorly about their weight (often thanks to public health campaigns), we actually have reason to suspect that the people who are not responding are more likely to be overweight than the people who do respond. Imputing a weight to Dan means that the number of overweight people in the sample will probably rise from 1 out of 3 (if we ignore Dan), to 2 out of 4 (if we impute Dan's weight). Clearly this matters. But doing it sensibly is more complicated than it sounds. Earlier, I suggested you should treat me like Carol, since we gave the same answer to the exercise question. But that's not quite right: there is a systematic difference between us. She answered the question, and I didn't. Given the social pressures faced by overweight people, isn't it likely that I'm *more* overweight than Carol? And of course this is still ignoring the fact that it's not sensible to impute a *single* weight to me, as if you actually knew my weight. Instead, what you need to do is impute a range of plausible guesses (referred to as multiple imputation), in order to capture the fact that you're more uncertain about my weight than you are about Carol's. And let's not get started on the problem posed by the fact that Elaine didn't send in the survey. As you can probably guess, dealing with missing data is an increasingly important topic. In fact, I've been told that a lot of journals in some fields will not accept studies that have missing data unless some kind of sensible multiple imputation scheme is followed.

- **Power analysis.** In Chapter 11 I discussed the concept of power (i.e., how likely are you to be able to detect an effect if it actually exists), and referred to power analysis, a collection of tools that are useful for assessing how much power your study has. Power analysis can be useful for planning a study (e.g., figuring out how large a sample you're likely to need), but it also serves a useful role in analysing data that you already collected. For instance, suppose you get a significant result, and you have an estimate of your effect size. You can use this information to estimate how much power your study actually had. This is kind of useful, especially if your effect size is not large. For instance, suppose you reject the null hypothesis $p < .05$, but you use power analysis to figure out that your estimated power was only .08. The significant result means that, if the null hypothesis was in fact true, there was a 5% chance of getting data like this. But the low power means that, even if the null hypothesis is false, the effect size was really as small as it looks, there was only an 8% chance of getting data like the one you did. This suggests that you need to be pretty cautious, because luck seems to have played a big part in your results, one way or the other!
- **Data analysis using theory-inspired models.** In a few places in this book I've mentioned response time (RT) data, where you record how long it takes someone to do something (e.g., make a simple decision). I've mentioned that RT data are almost invariably non-normal, and positively skewed. Additionally, there's a thing known as the speed-accuracy tradeoff: if you try to make decisions too quickly (low RT), you're likely to make poorer decisions (lower accuracy). So if you measure both the accuracy of a participant's decisions and their RT, you'll probably find that speed and accuracy are related. There's more to the story than this, of course, because some people make better decisions than others regardless of how fast they're going. Moreover, speed depends on both cognitive processes (i.e., time spent thinking) but also physiological ones (e.g., how fast can you move your muscles). It's starting to sound like analysing this data will be a complicated process. And indeed it is, but one of the things that you find when you dig into the psychological literature is that there already exist mathematical models (called "sequential sampling models")

that describe how people make simple decisions, and these models take into account a lot of the factors I mentioned above. You won't find any of these theoretically-inspired models in a standard statistics textbook. Standard stats textbooks describe standard tools, tools that could meaningfully be applied in lots of different disciplines, not just psychology. ANOVA is an example of a standard tool: it is just as applicable to psychology as to pharmacology. Sequential sampling models are not: they are psychology-specific, more or less. This doesn't make them less powerful tools: in fact, if you're analysing data where people have to make choices quickly, you should really be using sequential sampling models to analyse the data. Using ANOVA or regression or whatever won't work as well, because the theoretical assumptions that underpin them are not well-matched to your data. In contrast, sequential sampling models were explicitly designed to analyse this specific type of data, and their theoretical assumptions are *extremely* well-matched to the data. Obviously, it's impossible to cover this sort of thing properly, because there are thousands of context-specific models in every field of science. Even so, one thing that I'd like to do in later versions of the book is to give some case studies that are of particular relevance to psychologists, just to give a sense for how psychological theory can be used to do better statistical analysis of psychological data. So, in later versions of the book I'll probably talk about how to analyse response time data, among other things.

17.2

Learning the basics, and learning them in R

Okay, that was... long. And even that listing is massively incomplete. There really are a *lot* of big ideas in statistics that I haven't covered in this book. It can seem pretty depressing to finish a 500-page textbook only to be told that this is only the beginning, especially when you start to suspect that half of the stuff you've been taught is wrong. For instance, there are a lot of people in the field who would strongly argue against the use of the classical ANOVA model, yet I've devote two whole chapters to it! Standard ANOVA can be attacked from a Bayesian perspective, or from a robust statistics perspective, or even from a "it's just plain wrong" perspective (people very frequently use ANOVA when they should actually be using mixed models). So why learn it at all?

As I see it, there are two key arguments. Firstly, there's the pure pragmatism argument. Rightly or wrongly, ANOVA is widely used. If you want to understand the scientific literature, you need to understand ANOVA. And secondly, there's the "incremental knowledge" argument. In the same way that it was handy to have seen one-way ANOVA before trying to learn factorial ANOVA, understanding ANOVA is helpful for understanding more advanced tools, because a lot of those tools extend on or modify the basic ANOVA setup in some way. For instance, although mixed models are way more useful than ANOVA and regression, I've never heard of anyone learning how mixed models work without first having worked through ANOVA and regression. You have to learn to crawl before you can climb a mountain.

Actually, I want to push this point a bit further. One thing that I've done a lot of in this book is talk about fundamentals. I spent a lot of time on probability theory. I talked about the theory of estimation and hypothesis tests in more detail than I needed to. When talking about R, I spent a lot of time talking about how the language works, and talking about things like writing your own scripts, functions and programs. I didn't just teach you how to draw a histogram using `hist()`, I tried to give a basic overview of how the graphics system works. Why did I do all this? Looking back, you might ask whether I really *needed* to spend all that time talking about what a probability distribution is, or why there was even a section on probability density. If the goal of the book was to teach you how to run a *t*-test or an ANOVA, was all that really necessary? Or, come to think of it, why bother with R at all? There are lots of free alternatives out there: PSPP, for instance, is an SPSS-like clone that is totally free, has simple "point and click" menus, and can (I think) do every single analysis that I've talked about in this book. And you

can learn PSPP in about 5 minutes. Was this all just a huge waste of everyone's time???

The answer, I hope you'll agree, is no. The goal of an introductory stats is *not* to teach ANOVA. It's not to teach *t*-tests, or regressions, or histograms, or *p*-values. The goal is to start you on the path towards becoming a skilled data analyst. And in order for you to become a skilled data analyst, you need to be able to do more than ANOVA, more than *t*-tests, regressions and histograms. You need to be able to *think properly* about data. You need to be able to learn the more advanced statistical models that I talked about in the last section, and to understand the theory upon which they are based. And you need to have access to software that will let you use those advanced tools. And this is where – in my opinion at least – all that extra time I've spent on the fundamentals pays off. If you understand the graphics system in R, then you can draw the plots that *you* want, not just the canned plots that someone else has built into R for you. If you understand probability theory, you'll find it much easier to switch from frequentist analyses to Bayesian ones. If you understand the core mechanics of R, you'll find it much easier to generalise from linear regressions using `lm()` to using generalised linear models with `glm()` or linear mixed effects models using `lme()` and `lmer()`. You'll even find that a basic knowledge of R will go a long way towards teaching you how to use other statistical programming languages that are based on it. Bayesians frequently rely on tools like WinBUGS and JAGS, which have a number of similarities to R, and can in fact be called from within R. In fact, because R is the “lingua franca of statistics”, what you'll find is that most ideas in the statistics literature has been implemented somewhere as a package that you can download from CRAN. The same cannot be said for PSPP, or even SPSS.

In short, I think that the big payoff for learning statistics this way is *extensibility*. For a book that only covers the very basics of data analysis, this book has a massive overhead in terms of learning R, probability theory and so on. There's a whole lot of other things that it pushes you to learn besides the specific analyses that the book covers. So if your goal had been to learn how to run an ANOVA in the minimum possible time, well, this book wasn't a good choice. But as I say, I don't think that is your goal. I think you want to learn how to do data analysis. And if that really is your goal, you want to make sure that the skills you learn in your introductory stats class are naturally and cleanly extensible to the more complicated models that you need in real world data analysis. You want to make sure that you learn to use the same tools that real data analysts use, so that you can learn to do what they do. And so yeah, okay, you're a beginner right now (or you were when you started this book), but that doesn't mean you should be given a dumbed-down story, a story in which I don't tell you about probability density, or a story where I don't tell you about the nightmare that is factorial ANOVA with unbalanced designs. And it doesn't mean that you should be given baby toys instead of proper data analysis tools. Beginners aren't dumb; they just lack knowledge. What you need is *not* to have the complexities of real world data analysis hidden from you. What you need are the skills and tools that will let you handle those complexities when they inevitably ambush you in the real world.

And what I hope is that this book – or the finished book that this will one day turn into – is able to help you with that.

Author’s note – I’ve mentioned it before, but I’ll quickly mention it again. This reference list is appallingly incomplete. Please don’t assume that these are the only sources I’ve relied upon. The final version of this book will have a *lot* more references. And if you see anything clever sounding in this book that doesn’t seem to have a reference, I can absolutely promise you that the idea was someone else’s. This is an introductory textbook: none of the ideas are original. I’ll take responsibility for all the errors, but I can’t take credit for any of the good stuff. Everything smart in this book came from someone else, and they all deserve proper attribution for their excellent work. I just haven’t had the chance to give it to them yet.

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