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Second
Edition

Data Science at the Command Line

Obtain, Scrub, Explore, and Model Data
with Unix Power Tools



Jeroen Janssens
Foreword by Tim O'Reilly

Praise for *Data Science at the Command Line*

*Traditional computer and data science curricula all too often mistake the command line as an obsolete relic instead of teaching it as the modern and vital toolset that it is. Only well into my career did I come to grasp the elegance and power of the command line for easily exploring messy datasets and even creating reproducible data pipelines for work. The first edition of *Data Science at the Command Line* was one of the most comprehensive and clear references when I was a novice in the art, and now with the second edition, I'm again learning new tools and applications from it.*

—Dan Nguyen, data scientist, former news application developer at ProPublica, and former Lorry I. Lokey Visiting Professor in Professional Journalism at Stanford University

The Unix philosophy of simple tools, each doing one job well, then cleverly piped together, is embodied by the command line. Jeroen expertly discusses how to bring that philosophy into your work in data science, illustrating how the command line is not only the world of file input/output, but also the world of data manipulation, exploration, and even modeling.

—Chris H. Wiggins, associate professor in the department of applied physics and applied mathematics at Columbia University, and chief data scientist at The New York Times

This book explains how to integrate common data science tasks into a coherent workflow. It's not just about tactics for breaking down problems, it's also about strategies for assembling the pieces of the solution.

—John D. Cook, consultant in applied mathematics, statistics, and technical computing

Despite what you may hear, most practical data science is still focused on interesting visualizations and insights derived from flat files. Jeroen's book leans into this reality, and helps reduce complexity for data practitioners by showing how time-tested command-line tools can be repurposed for data science.

—Paige Bailey, principal product manager code intelligence at Microsoft, GitHub

It's amazing how fast so much data work can be performed at the command line before ever pulling the data into R, Python, or a database. Older technologies like sed and awk are still incredibly powerful and versatile. Until I read Data Science at the Command Line, I had only heard of these tools but never saw their full power. Thanks to Jeroen, it's like I now have a secret weapon for working with large data.

—Jared Lander, chief data scientist at Lander Analytics, organizer of the New York Open Statistical Programming Meetup, and author of R for Everyone

The command line is an essential tool in every data scientist's toolbox, and knowing it well makes it easy to translate questions you have of your data to real-time insights. Jeroen not only explains the basic Unix philosophy of how to chain together single-purpose tools to arrive at simple solutions for complex problems, but also introduces new command-line tools for data cleaning, analysis, visualization, and modeling.

—Jake Hofman, senior principal researcher at Microsoft Research, and adjunct assistant professor in the department of applied mathematics at Columbia University

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SECOND EDITION

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Jeroen Janssens

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by Jeroen Janssens

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[LSI]

Once again to my wife, Esther. Without her continued encouragement,
support,
and patience, this second edition would surely have ended up in /dev/null.

Foreword

It was love at first sight.

It must have been around 1981 or 1982 that I got my first taste of Unix. Its command-line shell, which uses the same language for single commands and complex programs, changed my world, and I never looked back.

I was a writer who had discovered the joys of computing, and regular expressions were my gateway drug. I'd first tried them in the text editor in HP's RTE operating system, but it was only when I came to Unix and its philosophy of small cooperating tools with the command-line shell as the glue that tied them together that I fully understood their power. Regular expressions in `ed`, `ex`, `vi` (now `vim`), and `emacs` were powerful, sure, but it wasn't until I saw how `ex` scripts unbound became `sed`, the Unix stream editor, and then AWK, which allowed you to bind programmed actions to regular expressions, and how shell scripts let you build pipelines not only out of the existing tools but out of new ones you'd written yourself, that I really got it. Programming is how you speak with computers, how you tell them what you want them to do, not just once, but in ways that persist, in ways that can be varied like human language, with repeatable structure but different verbs and objects.

As a beginner, other forms of programming seemed more like recipes to be followed exactly—careful incantations where you had to get everything right—or like waiting for a teacher to grade an essay you'd written. With shell programming, there was no compilation and waiting. It was more like a conversation with a friend. When the friend didn't understand, you could easily try again. What's more, if you had something simple to say, you could just say it with one word. And there were already words for a whole lot of the things you might want to say. But if there weren't, you could easily make up new words. And you could string together the words you learned and the words you made up into gradually more complex sentences, paragraphs, and eventually get to persuasive essays.

Almost every other programming language is more powerful than the shell and its associated tools, but for me at least, none provides an easier pathway

into the programming mindset, and none provides a better environment for a kind of everyday conversation with the machines that we ask to help us with our work. As Brian Kernighan, one of the creators of AWK as well as the coauthor of the marvelous book *The Unix Programming Environment*, said in an interview with Lex Fridman, “[Unix] was meant to be an environment where it was really easy to write programs.” [00:23:10] Kernighan went on to explain why he often still uses AWK rather than writing a Python program when he’s exploring data: “It doesn’t scale to big programs, but it does pretty darn well on these little things where you just want to see all the somethings in something.” [00:37:01]

In *Data Science at the Command Line*, Jeroen Janssens demonstrates just how powerful the Unix/Linux approach to the command line is even today. If Jeroen hadn’t already done so, I’d write an essay here about just why the command line is such a sweet and powerful match with the kinds of tasks so often encountered in data science. But he already starts out this book by explaining that. So I’ll just say this: the more you use the command line, the more often you will find yourself coming back to it as the easiest way to do much of your work. And whether you’re a shell newbie, or just someone who hasn’t thought much about what a great fit shell programming is for data science, this is a book you will come to treasure. Jeroen is a great teacher, and the material he covers is priceless.

Tim O'Reilly
May 2021

Preface

Data science is an exciting field to work in. It's also still relatively young. Unfortunately, many people, and many companies as well, believe that you need new technology to tackle the problems posed by data science. However, as this book demonstrates, many things can be accomplished by using the command line instead, and sometimes in a much more efficient way.

During my PhD program, I gradually switched from using Microsoft Windows to using Linux. Because this transition was a bit scary at first, I started with having both operating systems installed next to each other (known as a dual-boot). The urge to switch back and forth between Microsoft Windows and Linux eventually faded, and at some point I was even tinkering around with Arch Linux, which allows you to build up your own custom Linux machine from scratch. All you're given is the command line, and it's up to you what to make of it. Out of necessity, I quickly became very comfortable using the command line. Eventually, as spare time got more precious, I settled down with a Linux distribution known as Ubuntu because of its ease of use and large community. However, the command line is still where I'm spending most of my time.

It actually wasn't too long ago that I realized that the command line is not just for installing software, configuring systems, and searching files. I started learning about tools such as `cut`, `sort`, and `sed`. These are examples of command-line tools that take data as input, do something to it, and print the result. Ubuntu comes with quite a few of them. Once I understood the potential of combining these small tools, I was hooked.

After earning my PhD, when I became a data scientist, I wanted to use this approach to do data science as much as possible. Thanks to a couple of new, open source command-line tools including `xml2json`, `jq`, and `json2csv`, I was even able to use the command line for tasks such as scraping websites and processing lots of JSON data.

In September 2013, I decided to write a blog post titled “[7 Command-Line Tools for Data Science](#)”. To my surprise, the blog post got quite some attention, and I received a lot of suggestions of other command-line tools. I started wondering whether the blog post could be turned into a book. I was pleased that, some 10 months later, and with the help of many talented people (see the acknowledgments), the answer was yes.

I am sharing this personal story not so much because I think you should know how this book came about, but because I want to you know that I had to learn about the command line as well. Because the command line is so different from using a graphical user interface, it can seem scary at first. But if I could learn it, then you can as well. No matter what your current operating system is and no matter how you currently work with data, after reading this book you will be able to do data science at the command line. If you’re already familiar with the command line, or even if you’re already dreaming in shell scripts, chances are that you’ll still discover a few interesting tricks or command-line tools to use for your next data science project.

What to Expect from This Book

In this book, we’re going to obtain, scrub, explore, and model data—a lot of it. This book is not so much about how to become *better* at those data science tasks. There are already great resources available that discuss, for example, when to apply which statistical test or how data can best be visualized. Instead, this practical book aims to make you more *efficient* and *productive* by teaching you how to perform those data science tasks at the command line.

While this book discusses more than 90 command-line tools, it’s not the tools themselves that matter most. Some command-line tools have been around for a very long time, while others will be replaced by better ones. New command-line tools are being created even as you’re reading this. Over the years, I have discovered many amazing command-line tools.

Unfortunately, some of them were discovered too late to be included in the book. In short, command-line tools come and go. But that's OK.

What matters most is the underlying idea of working with tools, pipes, and data. Most command-line tools do one thing and do it well. This is part of the Unix philosophy, which makes several appearances throughout the book. Once you have become familiar with the command line, know how to combine command-line tools, and can even create new ones, you have developed an invaluable skill.

Changes for the Second Edition

While the command line as a technology and as a way of working is timeless, some of the tools discussed in the first edition have either been superseded by newer tools (e.g., `csvkit` has largely been replaced by `xsv`) or abandoned by their developers (e.g., `drake`), or they've been suboptimal choices (e.g., `weka`). I have learned a lot since the first edition was published in October 2014, either through my own experience or as a result of the useful feedback from my readers. Even though the book is quite niche because it lies at the intersection of two subjects, there remains a steady interest from the data science community, as evidenced by the many positive messages I receive almost every day. By updating the first edition, I hope to keep the book relevant for at least another five years. Here's a nonexhaustive list of changes I have made:

- I replaced `csvkit` with `xsv` as much as possible. `xsv` is a faster alternative to working with CSV files.
- In Chapters 2 and 3, I replaced the VirtualBox image with a Docker image. Docker is a faster and more lightweight way of running an isolated environment.
- I now use `pup` instead of `scrape` to work with HTML. `scrape` is a Python tool I created myself. `pup` is much faster, has more features, and is easier to install.

- Chapter 6 has been rewritten from scratch. Instead of `drake`, I now use `make` to do project management. `drake` is no longer maintained, and `make` is much more mature and very popular with developers.
- I replaced `Rio` with `rush`. `Rio` is a clunky Bash script I created myself. `rush` is an R package that is a much more stable and flexible way of using R from the command line.
- In Chapter 9 I replaced Weka and BigML with Vowpal Wabbit (`vw`). Weka is old, and the way it is used from the command line is clunky. BigML is a commercial API that I no longer want to rely on. Vowpal Wabbit is a very mature machine learning tool that was developed at Yahoo! and is now at Microsoft.
- Chapter 10 is an entirely new chapter about integrating the command line into existing workflows, including Python, R, and Apache Spark. In the first edition I mentioned that the command line can easily be integrated with existing workflows but never delved into the topic. This chapter fixes that.

How to Read This Book

In general, I advise you to read this book in a linear fashion. Once a concept or command-line tool has been introduced, chances are that I employ it in a later chapter. For example, in Chapter 9, I make heavy use of `parallel`, which is discussed extensively in Chapter 8.

Data science is a broad field that intersects many other fields such as programming, data visualization, and machine learning. As a result, this book touches on many interesting topics that unfortunately cannot be discussed at great length. At the end of each chapter, I provide suggestions for further exploration. It's not required that you read this material in order to follow along with the book, but if you are interested, just know that there's much more to learn.

Who This Book Is For

This book makes just one assumption about you: that you work with data. It doesn't matter which programming language or statistical computing environment you're currently using. The book explains all the necessary concepts from the beginning.

It also doesn't matter whether your operating system is Microsoft Windows, macOS, or some flavor of Linux. The book comes with a Docker image, which is an easy-to-install virtual environment. It allows you to run the command-line tools and follow along with the code examples in the same environment as this book was written. You don't have to waste time figuring out how to install all the command-line tools and their dependencies.

The book contains some code in Bash, Python, and R, so it's helpful if you have some programming experience, but it's by no means required to follow along with the examples.

Conventions Used in This Book

The following typographical conventions are used in this book:

Italic

Indicates new terms, URLs, directory names, and filenames.

Constant width

Used for code and commands, as well as within paragraphs to refer to command-line tools and their options.

Constant width bold

Shows commands or other text that should be typed literally by the user.

Constant width italic

Shows text that should be replaced with user-supplied values or by values determined by context.

TIP

This element signifies a tip or suggestion.

NOTE

This element signifies a general note.

WARNING

This element indicates a warning or caution.

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Acknowledgments for the Second Edition (2021)

Seven years have passed since the first edition came out. During this time, and especially during the last 13 months, many people have helped me. Without them, I would have never been able to write a second edition.

I was once again blessed with three wonderful editors at O'Reilly. I would like to thank Sarah "Embrace the deadline" Grey, Jess "Pedal to the metal" Haberman, and Kate "Let it go" Galloway. Their middle names say it all. With their incredible help, I was able to embrace the deadlines, put the pedal to metal when it mattered, and eventually let it go. I'd also like to thank their colleagues Angela Rufino, Arthur Johnson, Cassandra Furtado, David Futato, Helen Monroe, Karen Montgomery, Kate Dullea, Kristen Brown, Marie Beaugureau, Marsee Henon, Nick Adams, Regina Wilkinson, Shannon Cutt, Shannon Turlington, and Yasmina Greco, for making the collaboration with O'Reilly such a pleasure.

Despite having an automated process to execute the code and paste back the results (thanks to R Markdown and Docker), the number of mistakes I was able to make is impressive. Thank you Aaditya Maruthi, Brian Eoff, Caitlin Hudon, Julia Silge Mike Dewar, and Shane Reustle for reducing this number immensely. Of course, any mistakes left are my responsibility.

Marc Canaleta deserves a special thank you. In October 2014, shortly after the first edition came out, Marc invited me to give a one-day workshop about *Data Science at the Command Line* to his team at Social Point in Barcelona. Little did we both know that many workshops would follow. It eventually led me to start my own company: Data Science Workshops. Every time I teach, I learn something new. They probably don't know it, but each student has had an impact, in one way or another, on this book. To them I say: thank you. I hope I can teach for a very long time.

Captivating conversations, splendid suggestions, and passionate pull requests. I greatly appreciate each and every contribution by following generous people: Adam Johnson, Andre Manook, Andrea Borruso, Andres

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Throughout this book, and especially in the footnotes and appendix, you'll find hundreds of names. These names belong to the authors of the many tools, books, and other resources on which this book stands. I'm incredibly grateful for their hard work, regardless of whether that work was done 50 years or 50 days ago.

Above all, I would like to thank my wife Esther, my daughter Florien, and my son Olivier for reminding me daily what truly matters. I promise it'll be a few years before I start writing the third edition.

Acknowledgments for the First Edition (2014)

First of all, I'd like to thank Mike Dewar and Mike Loukides for believing that my blog post, "[7 Command-Line Tools for Data Science](#)", which I wrote in September 2013, could be expanded into a book.

Special thanks to my technical reviewers Mike Dewar, Brian Eoff, and Shane Reustle for reading various drafts, meticulously testing all the commands, and providing invaluable feedback. Your efforts have improved the book greatly. Any remaining errors are entirely my own responsibility.

I had the privilege of working with three amazing editors: Ann Spencer, Julie Steele, and Marie Beaugureau. Thank you for your guidance and for being such great liaisons with the many talented people at O'Reilly. Those people include Laura Baldwin, Huguette Barriere, Sophia DeMartini, Yasmina Greco, Rachel James, Ben Lorica, Mike Loukides, and Christopher Pappas. There are many others whom I haven't met because they are operating behind the scenes. Together they ensured that working with O'Reilly has truly been a pleasure.

This book discusses more than 80 command-line tools. Needless to say, without these tools, this book wouldn't have existed in the first place. I'm therefore extremely grateful to all the authors who created and contributed to these tools. The complete list of authors is unfortunately too long to include here; they are mentioned in the Appendix. Thanks especially to Aaron Crow, Jehiah Czebotar, Christoph Groskopf, Dima Kogan, Sergey Lisitsyn, Francisco J. Martin, and Ole Tange for providing help with their amazing command-line tools.

Eric Postma and Jaap van den Herik, who supervised me during my PhD program, deserve special thanks. Over the course of five years they taught me many lessons. Although writing a technical book is quite different from writing a PhD thesis, many of those lessons proved to be very helpful in the past nine months as well.

Finally, I'd like to thank my colleagues at YPlan, my friends, my family, and especially my wife, Esther, for supporting me and for pulling me away from the command line at just the right times.

Chapter 1. Introduction

This book is about doing data science at the command line. My aim is to make you a more efficient and productive data scientist by teaching you how to leverage the power of the command line.

Having both *data science* and *command line* in the book's title requires an explanation. How can a technology that is more than 50 years old¹ be of any use to a field that is only a few years young?

Today, data scientists can choose from an overwhelming collection of exciting technologies and programming languages. Python, R, Julia, and Apache Spark are but a few examples. You may already have experience in one or more of these. And if so, why should you still care about the command line for doing data science? What does the command line have to offer that these other technologies and programming languages do not?

These are valid questions. In this opening chapter I will answer these questions as follows. First, I provide a practical definition of data science that will act as the backbone of this book. Second, I'll list five important advantages of the command line. By the end of this chapter, I hope to have convinced you that the command line is indeed worth learning for doing data science.

Data Science Is OSEMN

The field of data science is still in its infancy, and as such, there exist various definitions of what it encompasses. Throughout this book I employ a very practical definition devised by Hilary Mason and Chris H. Wiggins.² They define data science according to the following five steps: (1) obtaining data, (2) scrubbing data, (3) exploring data, (4) modeling data, and (5) interpreting data. Together, these steps form the OSEMN (pronounced *awesome*) model. This definition serves as the backbone of this book

because each step (except for step 5, interpreting data, which I'll explain shortly) has its own chapter.

Although the five steps are discussed in a linear and incremental fashion, in practice it is very common to move back and forth between them or to perform multiple steps at the same time. **Figure 1-1** illustrates that doing data science is an iterative and non-linear process. For example, once you have modeled your data and have looked at the results, you may decide to go back to the scrubbing step to adjust the features of the dataset.

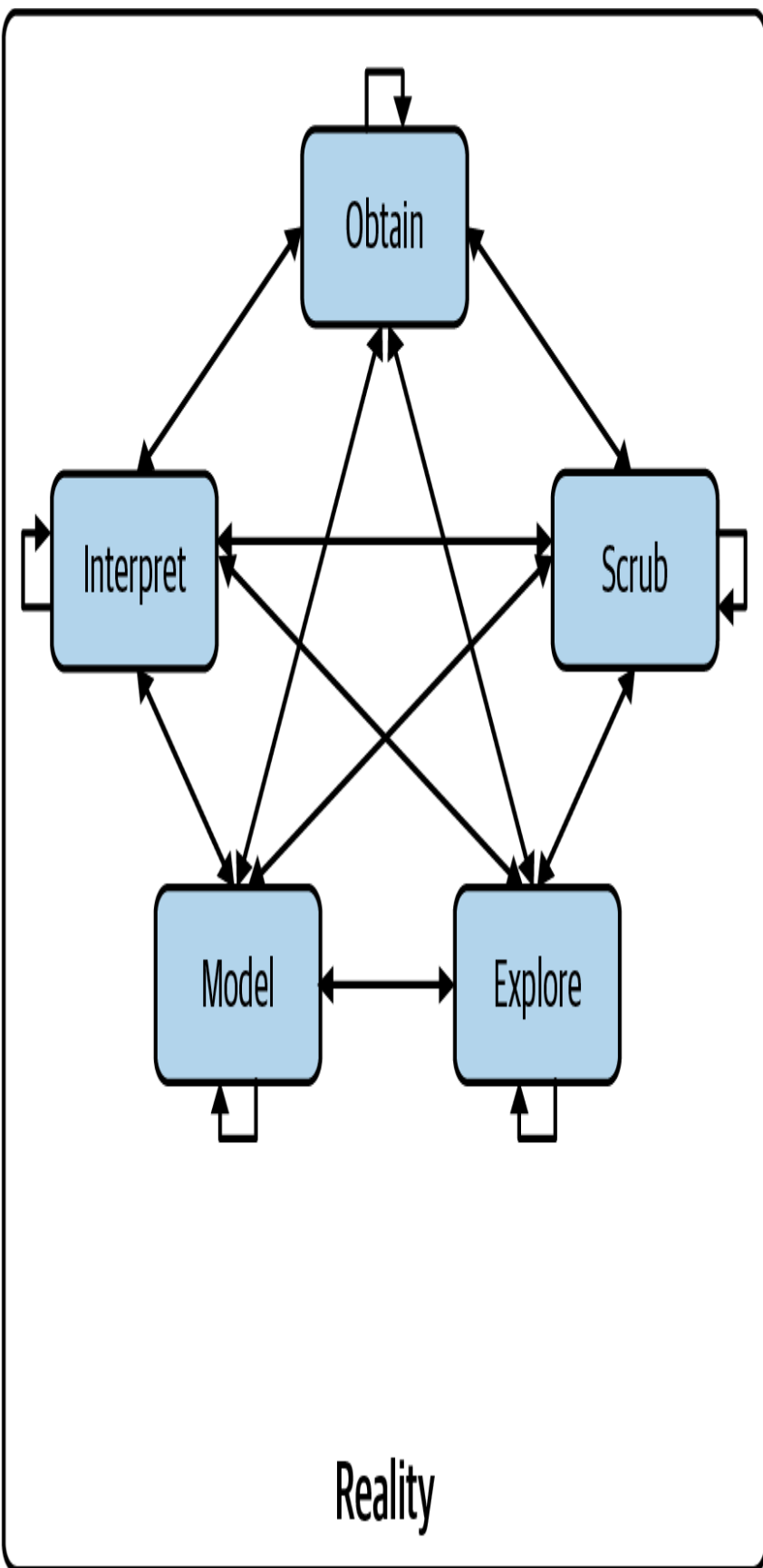
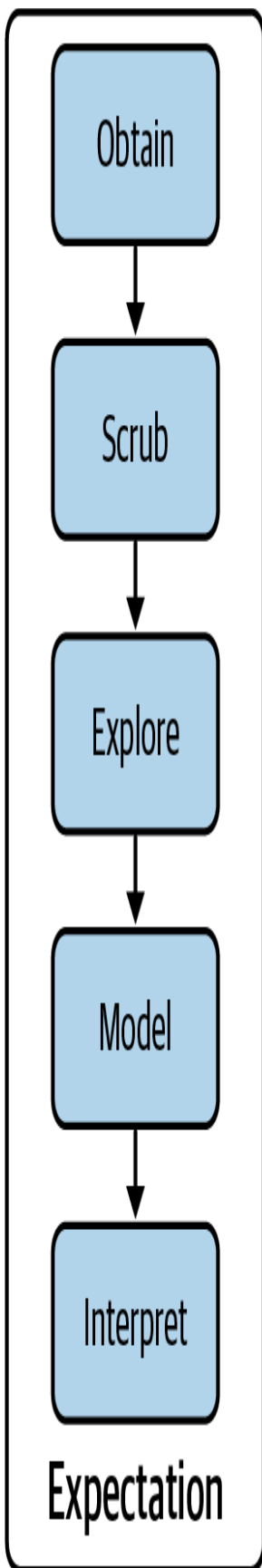


Figure 1-1. Doing data science is an iterative and nonlinear process

In the following pages, I explain what each step entails.

Obtaining Data

Without any data, there is little data science you can do. So the first step is obtaining data. Unless you are fortunate enough to already possess data, you may need to do one or more of the following:

- Download data from another location (e.g., a web page or server)
- Query data from a database or API (e.g., MySQL or Twitter)
- Extract data from another file (e.g., an HTML file or spreadsheet)
- Generate data yourself (e.g., reading sensors or taking surveys)

In [Chapter 3](#), I discuss several methods for obtaining data using the command line. The obtained data will most likely be in plain text, CSV, JSON, HTML, or XML format. The next step is to scrub this data.

Scrubbing Data

It is not uncommon for the obtained data to have missing values, inconsistencies, errors, weird characters, or uninteresting columns. In such cases, you have to *scrub*, or clean, the data before you can do anything interesting with it. Common scrubbing operations include:

- Filtering lines
- Extracting certain columns
- Replacing values
- Extracting words
- Handling missing values and duplicates
- Converting data from one format to another

While we data scientists love to create exciting data visualizations and insightful models (steps 3 and 4 of the OSEMN model), usually much effort goes into obtaining and scrubbing the required data first (steps 1 and 2). In *Data Jujitsu*(O'Reilly), DJ Patil states that “80% of the work in any data project is in cleaning the data.” In [Chapter 5](#), I demonstrate how the command line can help accomplish such data scrubbing operations.

Exploring Data

Once you have scrubbed your data, you are ready to explore it. This is where it gets interesting, because it's when you're exploring that you truly get to know your data. In [Chapter 7](#) I show you how the command line can be used to:

- Look at your data
- Derive statistics from your data
- Create insightful visualizations

Command-line tools used in [Chapter 7](#) include `csvstat` and `rush`.

Modeling Data

If you want to explain your data or predict what will happen, you probably want to create a statistical model of the data. Techniques to create a model include clustering, classification, regression, and dimensionality reduction. The command line is not suitable for programming a new type of model from scratch. It is, however, very useful to be able to build a model from the command line. In [Chapter 9](#) I will introduce several command-line tools that either build a model locally or employ an API to perform the computation in the cloud.

Interpreting Data

The final and perhaps most important step in the OSEMN model is interpreting data. This step involves:

- Drawing conclusions from your data
- Evaluating what your results mean
- Communicating your results

To be honest, the computer is of little use here, and the command line does not really come into play at this stage. Once you have reached this step, it's up to you. This is the only step in the OSEMN model that does not have its own chapter. Instead, I refer you to the book *Thinking with Data* by Max Shron (O'Reilly).

Intermezzo Chapters

Besides the chapters that cover the OSEMN steps, there are four intermezzo chapters. Each discusses a more general topic concerning data science and how the command line is employed for that. These topics are applicable to any step in the data science process.

In [Chapter 4](#), I discuss how to create reusable tools for the command line. These personal tools can come from long commands that you have typed on the command line or from existing code that you have written in, say, Python or R. Being able to create your own tools allows you to become more efficient and productive.

Because the command line is an interactive environment for doing data science, it can become challenging to keep track of your workflow. In [Chapter 6](#), I demonstrate a command-line tool called `make`, which allows you to define your data science workflow in terms of tasks and the dependencies between them. This tool increases the reproducibility of your workflow, not only for you but also for your colleagues and peers.

In [Chapter 8](#), I explain how your commands and tools can be sped up by running them in parallel. Using a command-line tool called GNU Parallel, you can apply command-line tools to very large datasets and run them on multiple cores or even on remote machines.

In [Chapter 10](#), I discuss how to employ the power of the command line in other environments and programming languages, such as R, RStudio, Python, Jupyter Notebooks, and even Apache Spark.

What Is the Command Line?

Before I discuss *why* you should use the command line for data science, let's take a peek at *what* the command line actually looks like (it may be already familiar to you). Figures [1-2](#) and [1-3](#) show a screenshot of the command line as it appears by default on macOS and Ubuntu, respectively. Ubuntu is a particular distribution of GNU/Linux, and it's the one I'll be using in this book.

```
com.docker.cli 7:22
```

```
$ whoami
dst
$ date
Thu 15 Apr 2021 11:04:56 AM CEST
$ echo 'The command line is awesome!' | cowsay -f tux
```

```
< The command line is awesome! >
```

```
-----
```

```
\ \
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 |:_/ |
 //   \\
 (|   | )
 /'\'_ _/\'\
 \___)=(_/_
```

```
$ |
```

Figure 1-2. Command line on macOS



jeroen@DESKTOP-B13E2TR: ~

```
$ whoami
jeroen
$ date
Thu Apr 15 12:04:25 CEST 2021
$ echo 'The command line is awesome!' | cowsay -f tux
```

```
< The command line is awesome! >
-----
      \
      \
      .--.
      |o_o |
      |:_/ |
      //   \ \
      (   |   )
      /' \_   _/ \
      \__)=(_/
```

\$

Figure 1-3. Command line on Ubuntu

The window shown in the two screenshots is called the *terminal*. This is the program that enables you to interact with the *shell*. It is the shell that executes the commands you type in. In [Chapter 2](#), I explain these two terms in more detail.

NOTE

I'm not showing the Microsoft Windows command line (also known as the Command Prompt or PowerShell), because it's fundamentally different from and incompatible with the commands presented in this book. The good news is that you can install a Docker image on Microsoft Windows so that you're able to follow along. Installation of the Docker image is explained in [Chapter 2](#).

Interacting with your computer by typing commands is very different from going through a *graphical user interface* (GUI). If you are mostly used to processing data in, say, Microsoft Excel, then this approach may seem intimidating at first. Don't be afraid. Trust me when I say that you'll get used to working at the command line very quickly.

In this book, the commands that I type and the output that they generate are displayed as text. For example, the contents of the terminal in the two screenshots would look like this:

```
$ whoami  
dst  
  
$ date  
Tue Jun 29 02:25:17 PM CEST 2021  
  
$ echo 'The command line is awesome!' | cowsay -f tux  
< The command line is awesome! >  
-----  
 \ /  
  ...  
 |o_o |  
 |:_/ |  
 // \ \ )  
(| | )
```

'__ -/____)=(__/'

\$

You'll notice that each command is preceded by a dollar sign (`$`). This is called the *prompt*. The prompt in the two screenshots shows more information, namely the username, the date, and a penguin. It's a convention to show only a dollar sign in examples, because the prompt (1) can change during a session (when you go to a different directory), (2) can be customized by the user (e.g., it can also show the time or the current git³ branch you're working on), and (3) is irrelevant for the commands themselves.

In the next chapter I'll explain much more about essential command-line concepts. But first, it's time to explain *why* you should learn to use the command line for doing data science.

Why Data Science at the Command Line?

The command line has many great advantages that can really make you a more efficient and productive data scientist. Roughly grouping the advantages, the command line is *agile*, *augmenting*, *scalable*, *extensible*, and *ubiquitous*.

The Command Line Is Agile

The first advantage of the command line is that it allows you to be agile. Data science has a very interactive and exploratory nature, and the environment that you work in needs to allow for that. The command line achieves this by two means.

First, the command line provides a so-called *read-eval-print loop* (REPL). This means that you type in a command, press Enter, and the command is evaluated immediately. A REPL is often much more convenient for doing data science than the edit-compile-run-debug cycle associated with scripts,

large programs, and, say, Hadoop jobs. Your commands are executed immediately, may be stopped at will, and can be changed quickly. This short iteration cycle really allows you to play with your data.

Second, the command line is very close to the filesystem. Because data is the main ingredient for doing data science, it is important to be able to work easily with the files that contain your dataset. The command line offers many convenient tools for this.

The Command Line Is Augmenting

The command line integrates well with other technologies. Whatever technology your data science workflow currently includes (whether it's R, Python, or Excel), please know that I'm not suggesting you abandon that workflow. Instead, consider the command line as an augmenting technology that amplifies the technologies you're currently employing. It can do so in three ways.

First, the command line can act as a glue between many different data science tools. One way to glue tools is by connecting the output from the first tool to the input of the second tool. In [Chapter 2](#) I explain how this works.

Second, you can often delegate tasks to the command line from your own environment. For example, Python, R, and Apache Spark allow you to run command-line tools and capture their output. I demonstrate this with examples in [Chapter 10](#).

Third, you can convert your code (e.g., a Python or R script) into a reusable command-line tool. That way, the language that it's written in doesn't matter anymore; it can be used from the command line directly or from any environment that integrates with the command line, as mentioned in the previous paragraph. I explain how to do this in [Chapter 4](#).

In the end, every technology has its strengths and weaknesses, so it's good to know several technologies and use the one that is most appropriate for the task at hand. Sometimes that means using R, sometimes the command

line, and sometimes even pen and paper. By the end of this book you'll have a solid understanding of when you should use the command line, and when you're better off continuing with your favorite programming language or statistical computing environment.

The Command Line Is Scalable

As I've said before, working on the command line is very different from using a GUI. On the command line you do things by typing, whereas with a GUI you do things by pointing and clicking with a mouse.

Everything that you type manually on the command line can also be automated through scripts and tools. This makes it very easy to rerun your commands if you made a mistake, when the input data has changed, or because your colleague wants to perform the same analysis. Moreover, your commands can be run at specific intervals, on a remote server, and in parallel on many chunks of data (more on that in [Chapter 8](#)).

Because the command line is automatable, it becomes scalable and repeatable. It's not straightforward to automate pointing and clicking, which makes a GUI a less suitable environment for doing scalable and repeatable data science.

The Command Line Is Extensible

The command line itself was invented over 50 years ago. Its core functionality has largely remained unchanged, but its *tools*, which are the workhorses of the command line, are being developed on a daily basis.

The command line itself is language agnostic. This allows the command-line tools to be written in many different programming languages. The open source community is producing many free and high-quality command-line tools that we can use for data science.

These command-line tools can work together, which makes the command line very flexible. You can also create your own tools, allowing you to extend the effective functionality of the command line.

The Command Line Is Ubiquitous

Because the command line comes with any Unix-like operating system, including Ubuntu Linux and macOS, it can be found in many places. Plus, 100% of the top five hundred supercomputers are running Linux.⁴ So if you ever get your hands on one of those supercomputers (or if you ever find yourself in Jurassic Park with the door locks not working), you'd better know your way around the command line!

But Linux doesn't run only on supercomputers. It also runs on servers, laptops, and embedded systems. These days, many companies offer cloud computing, where you can easily launch new machines on the fly. If you ever log in to such a machine (or a server in general), it's almost certain that you'll arrive at the command line.

It's also important to note that the command line isn't just hype. This technology has been around for more than five decades, and I'm convinced that it's here to stay for another five. Learning how to use the command line (for data science and in general) is therefore a worthwhile investment.

Summary

In this chapter I have introduced you to the OSEMN model for doing data science, which I use as a guide throughout the book. I have provided some background about the Unix command line and hopefully convinced you that it's a suitable environment for doing data science. In the next chapter I'll show you how to get started by installing the datasets and tools and explain the fundamental concepts.

For Further Exploration

- The book *UNIX: A History and a Memoir* by Brian W. Kernighan (self-published) tells the story of Unix, explaining what it is, how it was developed, and why it matters.

- In 2018, I gave a presentation titled “50 Reasons to Learn the Shell for Doing Data Science” at Strata London. You can [read the slides](#) if you need even more convincing.
- The short but sweet book *Thinking with Data* by Max Shron ([O'Reilly](#)) focuses on the *why* instead of the *how* and provides a framework for defining your data science project that will help you ask the right questions and solve the right problems.

-
- 1 The development of the UNIX operating system started back in 1969. It featured a command line since the beginning. The important concept of pipes, which I will discuss in “[Essential Unix Concepts](#)”, was added in 1973.
 - 2 “A Taxonomy of Data Science,” *dataists* (blog), September 25, 2010, <http://www.dataists.com/2010/09/a-taxonomy-of-data-science>.
 - 3 Linus Torvalds and Junio C. Hamano, *git – the Stupid Content Tracker*, version 2.25.1, 2021, <https://git-scm.com>.
 - 4 See [TOP500](#), which keeps track of how many supercomputers run Linux.

Chapter 2. Getting Started

In this chapter, I'm going to make sure that you have all the prerequisites for doing data science at the command line. The prerequisites are threefold: (1) having the same datasets that I use in this book, (2) having a proper environment with all the command-line tools that I use throughout this book, and (3) understanding the essential concepts that come into play when using the command line.

First, I describe how to download the datasets. Second, I explain how to install the Docker image, which is a virtual environment based on Ubuntu Linux that contains all the necessary command-line tools. Finally, I go over the essential Unix concepts through examples.

By the end of this chapter, you'll have everything you need to continue with the first step of doing data science, namely obtaining data.

Getting the Data

The datasets I use in this book can be obtained as follows:

1. Download the ZIP file from [the book's website](#).
2. Create a new directory. You can give this directory any name you like, but I recommend you stick to lowercase letters, numbers, and maybe a hyphen or an underscore so that the name is easier to work with at the command line—for example, *dsatcl2*. Remember where this directory is.
3. Move the ZIP file to that new directory and unpack it.
4. This directory now contains one subdirectory per chapter.

In the next section I explain how to install the environment containing all the command-line tools to work with this data.

Installing the Docker Image

In this book we use many different command-line tools. Unix often comes with a lot of command-line tools preinstalled and offers many packages that contain more relevant tools. Installing these packages yourself is often not too difficult. However, we'll also use tools that are not available as packages and require a more manual and more involved installation. So that you can acquire the necessary command-line tools without having to go through the installation process for each tool, I encourage you, whether you're on Windows, macOS, or Linux, to install the Docker image that was created specifically for this book.

A Docker image is a bundle of one or more applications together with all their dependencies. A Docker container is an isolated environment that runs an image. You can manage Docker images and containers using the `docker` command-line tool (which is what you'll do below) or the Docker GUI. In a way, a Docker container is like a virtual machine, only a Docker container uses far fewer resources. At the end of this chapter I suggest some resources for learning more about Docker.

TIP

If you still prefer to run the command-line tools natively rather than inside a Docker container, then you can, of course, install the command-line tools individually yourself. The code to build the Docker image can be found [on GitHub](#) and may serve as a guide to help you with that. Please be aware that this can be time consuming for some tools, as they require many nontrivial steps, such as compiling from source.

To install the Docker image, you first need to download Docker itself from [the Docker website](#). Once it is installed, you invoke the following command on your terminal or command prompt to download the Docker image (don't type the dollar sign):

```
$ docker pull datasciencetoolbox/dsatcl2e
```

You can run the Docker image as follows:

```
$ docker run --rm -it datasciencetoolbox/dsatcl2e
```

You're now inside an isolated environment known as a *Docker container* that has all the necessary command-line tools installed. If the following command produces an enthusiastic cow, then you know everything is working correctly:

```
$ cowsay "Let's moove!"
```

```
< Let's moove! >
-----
 \  ^__^
  \  (oo)\_____
    (__)\       )\/\
        ||----w |
        ||     ||
```

If you want to get data in and out of the container, you can add a volume, which means that a local directory gets mapped to a directory inside the container. I recommend that you first create a new directory, navigate to this new directory, and then run the following when you're on macOS or Linux:

```
$ docker run --rm -it -v "$(pwd)":/data datasciencetoolbox/dsatcl2e
```

Or run the following when you're on Windows and using the Command Prompt (also known as `cmd`):

```
C:\> docker run --rm -it -v "%cd%":/data datasciencetoolbox/dsatcl2e
```

Or the following when you're using Windows PowerShell:

```
PS C:\> docker run --rm -it -v ${PWD}:data datasciencetoolbox/dsatcl2e
```

In the above commands, the option `-v` instructs `docker` to map the current directory to the `/data` directory inside the container, so this is the place to get data in and out of the Docker container.

NOTE

If you would like to know more about the Docker image, you can [visit it on Docker Hub](#).

When you’re done, you can shut down the Docker container by typing `exit`.

Essential Unix Concepts

In [Chapter 1](#), I briefly showed you what the command line is. Now that you are running the Docker image, we can really get started. In this section, I discuss several concepts and tools that you will need to know to feel comfortable doing data science at the command line. If up until now you have been mainly working with graphical user interfaces, then this might be quite a change. But don’t worry—I’ll start at the beginning and very gradually go on to more advanced topics.

NOTE

This section is not a complete course in Unix. I will explain only the concepts and tools that are relevant to doing data science. One of the advantages of the Docker image is that a lot is already set up. If you wish to know more, consult “[For Further Exploration](#)”.

The Environment

So you’ve just logged in to a brand-new environment. Before you do anything, it’s worthwhile to get a high-level understanding of this environment, which is roughly defined by four layers, listed here from the top down:

Command-line tools

First and foremost, there are the command-line tools that you work with. We use them by typing their corresponding commands. There are

different types of command-line tools, which I will discuss in the next section. Examples of tools are `ls`,¹ `cat`,² and `jq`.³

Terminal

The terminal, which is the second layer, is the application that we type our commands in. If you see the following text mentioned in the book:

```
$ seq 3  
1  
2  
3
```

then you would type `seq 3` into your terminal and press Enter. (The command-line tool `seq`,⁴ as you can see, generates a sequence of numbers.) You do not type the dollar sign (\$). It's just there to tell you that this is a command you can type in the terminal. This dollar sign is known as the *prompt*. The text below `seq 3` is the output of the command.

Shell

The third layer is the shell. Once we have typed in our command and pressed Enter, the terminal sends that command to the shell. The shell is a program that interprets the command. I use the Z shell, but many other shells are available, such as Bash and Fish.

Operating system

The fourth layer is the operating system, which is GNU/Linux in our case. Linux is the name of the kernel, which is the heart of the operating system. The kernel is in direct contact with the CPU, disks, and other hardware. The kernel also executes our command-line tools. GNU, which stands for “GNU’s not UNIX,” refers to the set of basic tools.

The Docker image is based on a particular GNU/Linux distribution called Ubuntu.

Executing a Command-Line Tool

Now that you have a basic understanding of the environment, it is high time that you try out some commands. Type the following in your terminal (without the dollar sign) and press Enter:

```
$ pwd  
/home/dst
```

You just executed a command that contained a single command-line tool. The tool `pwd`⁵ outputs the name of the directory where you currently are. By default, when you log in, this is your home directory.

The command-line tool `cd`, which is a Z shell builtin, allows you to navigate to a different directory:

```
$ cd /data/ch02 ①
```

```
$ pwd ②  
/data/ch02
```

```
$ cd .. ③
```

```
$ pwd ④  
/data
```

```
$ cd ch02 ⑤
```

- ① Navigate to the directory `/data/ch02`.
- ② Print the current directory.
- ③ Navigate to the parent directory.
- ④

Print the current directory again.

- ⑤ Navigate to the subdirectory `ch02`.

The part after `cd` specifies the directory you want to navigate to. Values that come after the command are called *command-line arguments* or *options*.

The two dots refer to the parent directory. One dot, by the way, refers to the current directory. While `cd .` wouldn't have any effect, you'll still see one dot being used in other places.

Let's try a different command:

```
$ head -n 3 movies.txt
Matrix
Star Wars
Home Alone
```

Here we pass three command-line arguments to `head`.⁶ The first one is an option. Here I used the short option `-n`. Sometimes a short option has a long variant, which would be `--lines` in this case. The second one is a value that belongs to the option. The third one is a filename. This particular command outputs the first three lines of the file `/data/ch02/movies.txt`.

Five Types of Command-Line Tools

I use the term *command-line tool* a lot, but I haven't yet explained what I actually mean by it. I use command-line tool as an umbrella term for *anything* that can be executed from the command line (see [Figure 2-1](#)). Under the hood, each command-line tool is one of the following five types:

- A binary executable
- A shell builtin
- An interpreted script
- A shell function

- An alias

Command-line tool

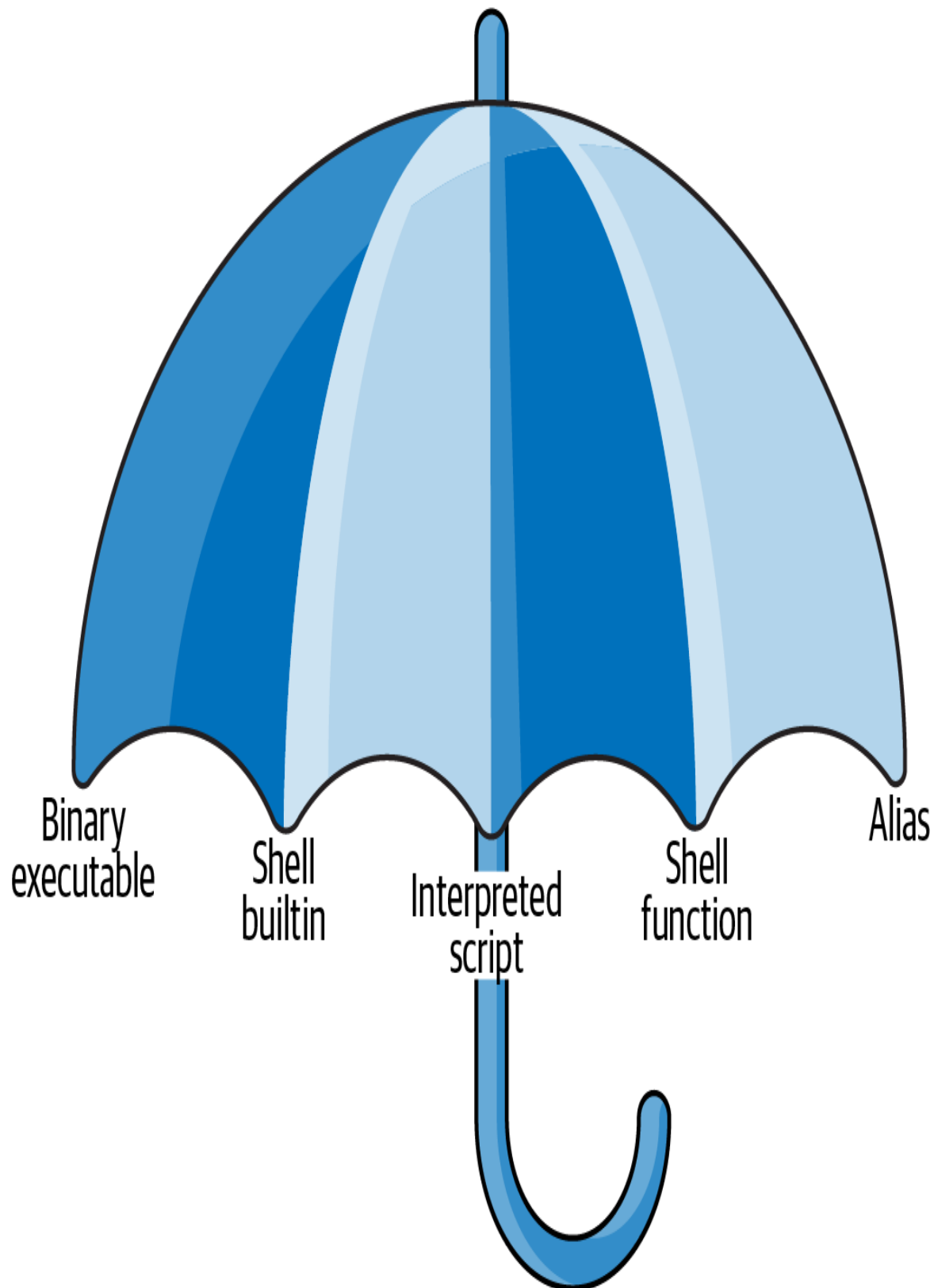


Figure 2-1. I use the term “command-line tool” as an umbrella term

It’s good to know the difference between the types. The command-line tools that come preinstalled with the Docker image mostly comprise the first two types (binary executable and shell builtin). The other three types (interpreted script, shell function, and alias) allow us to further build up our data science toolbox and become more efficient and more productive data scientists:

Binary executable

Binary executables are programs in the classical sense. A binary executable is created by compiling source code to machine code. This means that when you open the file in a text editor, you cannot read it.

Shell builtin

Shell builtins are command-line tools provided by the shell, which is the Z shell (or zsh) in our case. Examples include `cd` and `pwd`. Shell builtins may differ between shells. Like binary executables, they cannot be easily inspected or changed.

Interpreted script

An interpreted script is a text file that is executed by a binary executable. Examples include Python, R, and Bash scripts. One great advantage of an interpreted script is that you can read and change it. The following script is interpreted by Python not because of the file extension `.py` but because the first line of the script defines the binary that should execute it:

```
$ bat fac.py
-----
| File: fac.py
-----
1 | #!/usr/bin/env python
```

```
2 |
3 | def factorial(x):
4 |     result = 1
5 |     for i in range(2, x + 1):
6 |         result *= i
7 |     return result
8 |
9 | if __name__ == "__main__":
10 |     import sys
11 |     x = int(sys.argv[1])
12 |     sys.stdout.write(f"{factorial(x)}\n")
```

This script computes the factorial of the integer that we pass as a parameter. It can be invoked from the command line as follows:

```
$ ./fac.py 5
```

120

In [Chapter 4](#), I'll discuss in great detail how to create reusable command-line tools using interpreted scripts.

Shell function

A shell function is a function that is executed by the shell itself (zsh, in our case). Shell functions provide similar functionality to a script, but they are usually (but not necessarily) smaller than scripts. They also tend to be more personal. The following command defines a function called `fac`, which, just like the interpreted Python script I just described, computes the factorial of the integer we pass as a parameter. It does so by generating a list of numbers using `seq`, putting those numbers on one

line with * as the delimiter using `paste`,⁷ and passing this equation into `bc`,⁸ which evaluates it and outputs the result:

```
$ fac() { (echo 1; seq $1) | paste -s -d\* - | bc; }
```

```
$ fac 5
```

```
120
```

The file `~/.zshrc`, which is a configuration file for the Z shell, is a good place to define your shell functions so that they are always available.

Alias

Aliases are like macros. If you often find yourself executing a certain command with some or all of the same parameters, you can define an alias for the command to save time. An alias is also very useful when you continue to misspell a certain command (Chris Wiggins maintains a [useful list of aliases](#)). The following command defines such an alias:

```
$ alias l='ls --color -lhF --group-directories-first'
```

```
$ alias less=less
```

Now, if you type the following on the command line, the shell will replace each alias it finds with its value:

```
$ cd /data
```

```
$ l
```

```
total 40K
```

```
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch01/
```

```
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch02/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch03/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch04/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch05/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch06/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch07/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch08/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch09/
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch10/
```

```
$ cd ch02
```

Aliases are simpler than shell functions, as they don't allow parameters. The function `fac` could not have been defined using an alias because of the parameter. Still, aliases allow you to save lots of keystrokes. Like shell functions, aliases are often defined in the file `.zshrc`, which is located in your home directory. To see all aliases currently defined, you run `alias` without arguments. Try it. What do you see?

In this book I focus mostly on the last three types of command-line tools: interpreted scripts, shell functions, and aliases. I do so because these tools can easily be changed. The purpose of a command-line tool is to make your life easier and to make you a more productive and more efficient data scientist. You can find out the type of a command-line tool with `type` (which is itself a shell builtin):

```
$ type -a pwd
pwd is a shell builtin
pwd is /usr/bin/pwd
pwd is /bin/pwd
```

```
$ type -a cd
```

```
cd is a shell builtin

$ type -a fac
fac is a shell function

$ type -a l
l is an alias for ls --color -lhF --group-directories-first
```

`type` returns three command-line tools for `pwd`. In that case, the first reported command-line tool is used when you type `pwd`. In the next section we'll look at how to combine command-line tools.

Combining Command-Line Tools

Because most command-line tools adhere to the Unix philosophy,⁹ they are designed to do only one thing, and to do it really well. For example, the command-line tool `grep`¹⁰ can filter lines, `wc`¹¹ can count lines, and `sort`¹² can sort lines. The power of the command line comes from its ability to combine these small yet powerful command-line tools.

This power is made possible by managing the communication streams of these tools. Each tool has three standard communication streams: standard input, standard output, and standard error. These are often abbreviated as `stdin`, `stdout`, and `stderr`.

Both the standard output and standard error are, by default, redirected to the terminal, so that both normal output and any error messages are printed on the screen. [Figure 2-2](#) illustrates this for both `pwd` and `rev`.¹³ If you run `rev`, you'll see that nothing happens. That's because `rev` expects input, which by default is any keys pressed on the keyboard. Try typing a sentence and pressing Enter—`rev` immediately responds with your input in reverse. You can stop sending input by pressing Ctrl-D after which `rev` will stop.

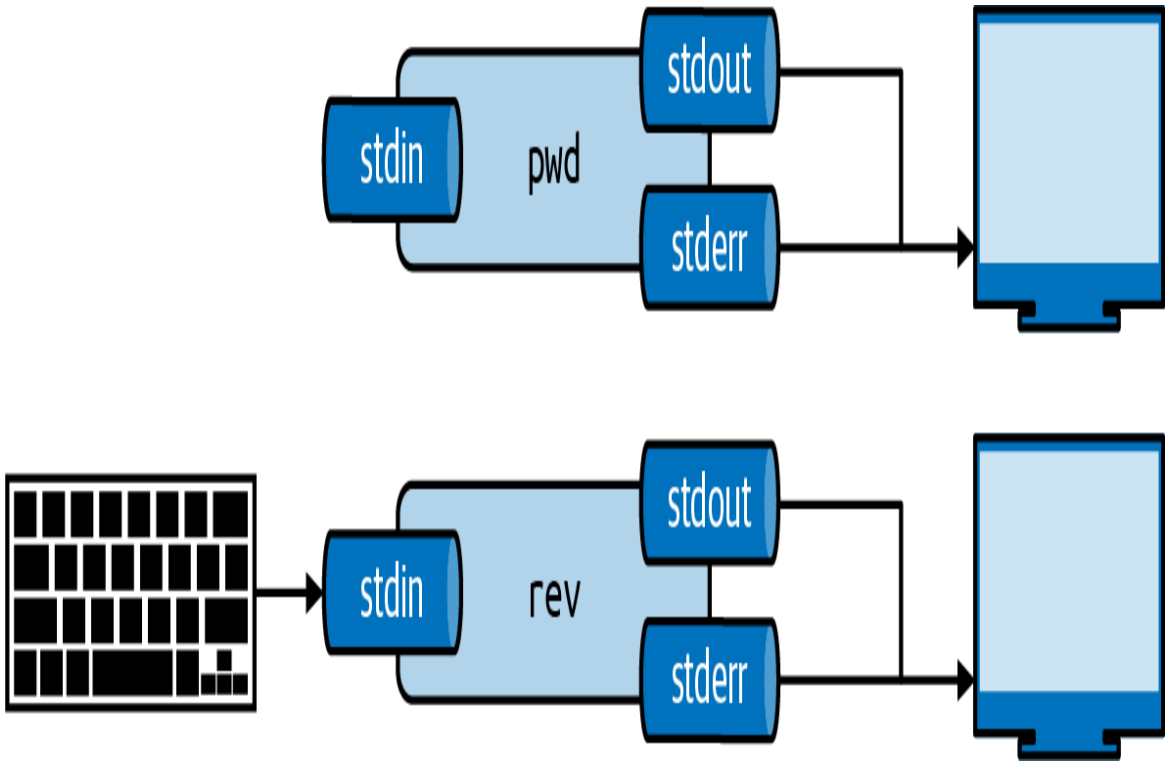


Figure 2-2. Every tool has three standard streams: standard input (`stdin`), standard output (`stdout`), and standard error (`stderr`)

In practice, rather than using the keyboard as a source of input, you'll use the output generated by other tools and the contents of files. For example, with `curl` we can download the book *Alice's Adventures in Wonderland* by Lewis Carroll and *pipe* that to the next tool; [Figure 2-3](#) illustrates piping the output from one tool to another tool. (I'll discuss `curl` in more detail in [Chapter 3](#).) This is done using the pipe operator (`|`).

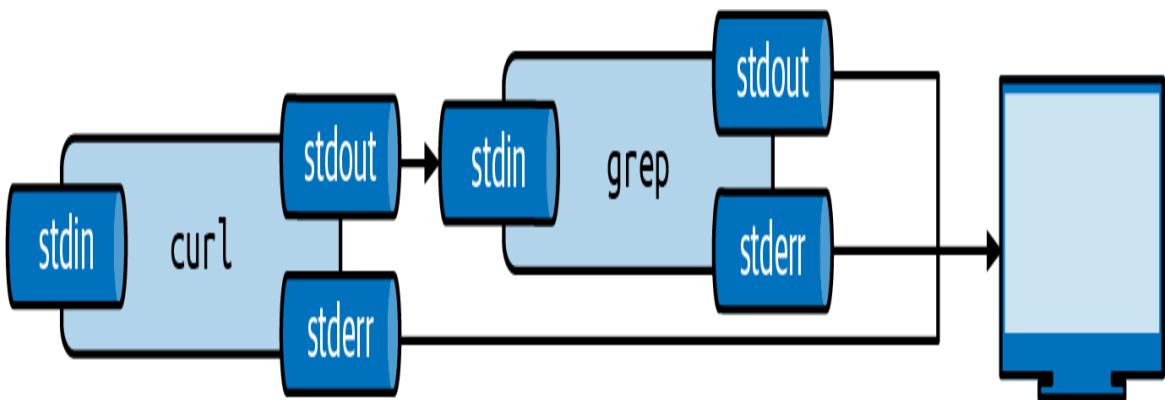


Figure 2-3. The output from a tool can be piped to another tool

We can pipe the output of `curl` to `grep` to filter lines on a pattern. Imagine that we want to see the chapters listed in the table of contents—we can combine `curl` and `grep` as follows:

```
$ curl -s "https://www.gutenberg.org/files/11/11-0.txt" | grep " CHAPTER"
CHAPTER I.      Down the Rabbit-Hole
CHAPTER II.     The Pool of Tears
CHAPTER III.    A Caucus-Race and a Long Tale
CHAPTER IV.     The Rabbit Sends in a Little Bill
CHAPTER V.      Advice from a Caterpillar
CHAPTER VI.     Pig and Pepper
CHAPTER VII.    A Mad Tea-Party
CHAPTER VIII.   The Queen's Croquet-Ground
CHAPTER IX.     The Mock Turtle's Story
CHAPTER X.      The Lobster Quadrille
CHAPTER XI.    Who Stole the Tarts?
CHAPTER XII.   Alice's Evidence
```

And if we want to know *how many* chapters the book has, we can use `wc`, which is very good at counting things:

```
$ curl -s "https://www.gutenberg.org/files/11/11-0.txt" |
> grep " CHAPTER" |
> wc -l ❶
12
```

- ❶ The option `-l` specifies that `wc` should output only the number of lines that are passed into it. By default, it also returns the number of characters and words.

You can think of piping as an automated copy and paste. Once you get the hang of combining tools using the pipe operator, you'll find that there are virtually no limits to the combinations you can make.

Redirecting Input and Output

Besides piping the output from one tool to another tool, you can also save it to a file. The file will be saved in the current directory, unless a full path is

given. This is called *output redirection*, and it works as follows:

```
$ curl "https://www.gutenberg.org/files/11/11-0.txt" | grep " CHAPTER" > chapter  
s.txt  
          % Total    % Received % Xferd  Average Speed   Time     Time     Time  
Current                                         Dload  Upload   Total   Spent   Left  Speed  
100  170k  100  170k    0      0  231k      0 --:--:-- --:--:-- --:--:--  231k  
  
$ cat chapters.txt  
CHAPTER I.      Down the Rabbit-Hole  
CHAPTER II.     The Pool of Tears  
CHAPTER III.    A Caucus-Race and a Long Tale  
CHAPTER IV.     The Rabbit Sends in a Little Bill  
CHAPTER V.      Advice from a Caterpillar  
CHAPTER VI.     Pig and Pepper  
CHAPTER VII.    A Mad Tea-Party  
CHAPTER VIII.   The Queen's Croquet-Ground  
CHAPTER IX.     The Mock Turtle's Story  
CHAPTER X.      The Lobster Quadrille  
CHAPTER XI.    Who Stole the Tarts?  
CHAPTER XII.   Alice's Evidence
```

Here, we save the output of `grep` to a file named `chapters.txt` in the directory `/data/ch02`. If this file does not exist yet, it will be created. If this file already exists, its contents are overwritten. [Figure 2-4](#) illustrates how output redirection works conceptually. Note that the standard error is still redirected to the terminal.

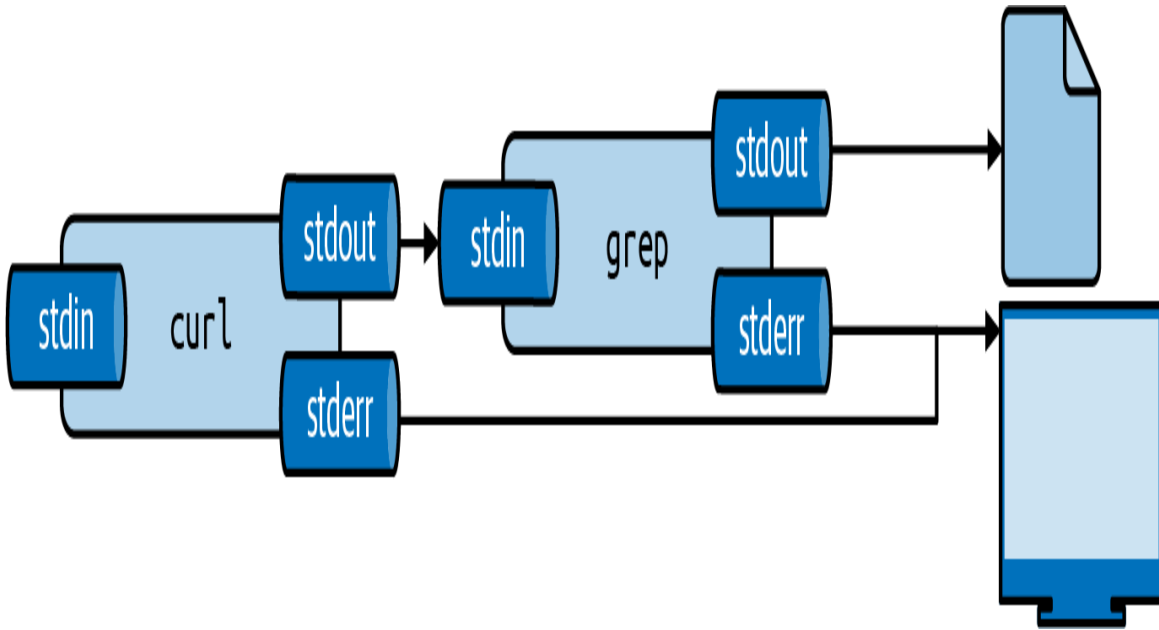


Figure 2-4. The output from a tool can be redirected to a file

You can also append the output to a file with `>>`, meaning the output is added after the original contents:

```
$ echo -n "Hello" > greeting.txt
$ echo " World" >> greeting.txt
```

The tool `echo` outputs the value you specify. The `-n` option, which stands for *newline*, specifies that `echo` should not output a trailing newline.

Saving the output to a file is useful if you need to store intermediate results, for example, to continue with your analysis at a later stage. To use the contents of the file `greeting.txt` again, we can use `cat`, which reads a file and prints it:

```
$ cat greeting.txt
Hello World

$ cat greeting.txt | wc -w ①
2
```

① The `-w` option instructs `wc` to count only words.

The same result can be achieved by using the less-than sign (<):

```
$ < greeting.txt wc -w  
2
```

This way, you are directly passing the file to the standard input of `wc` without running an additional process.¹⁴ Figure 2-5 illustrates how these two ways work. Again, the final output is the same.

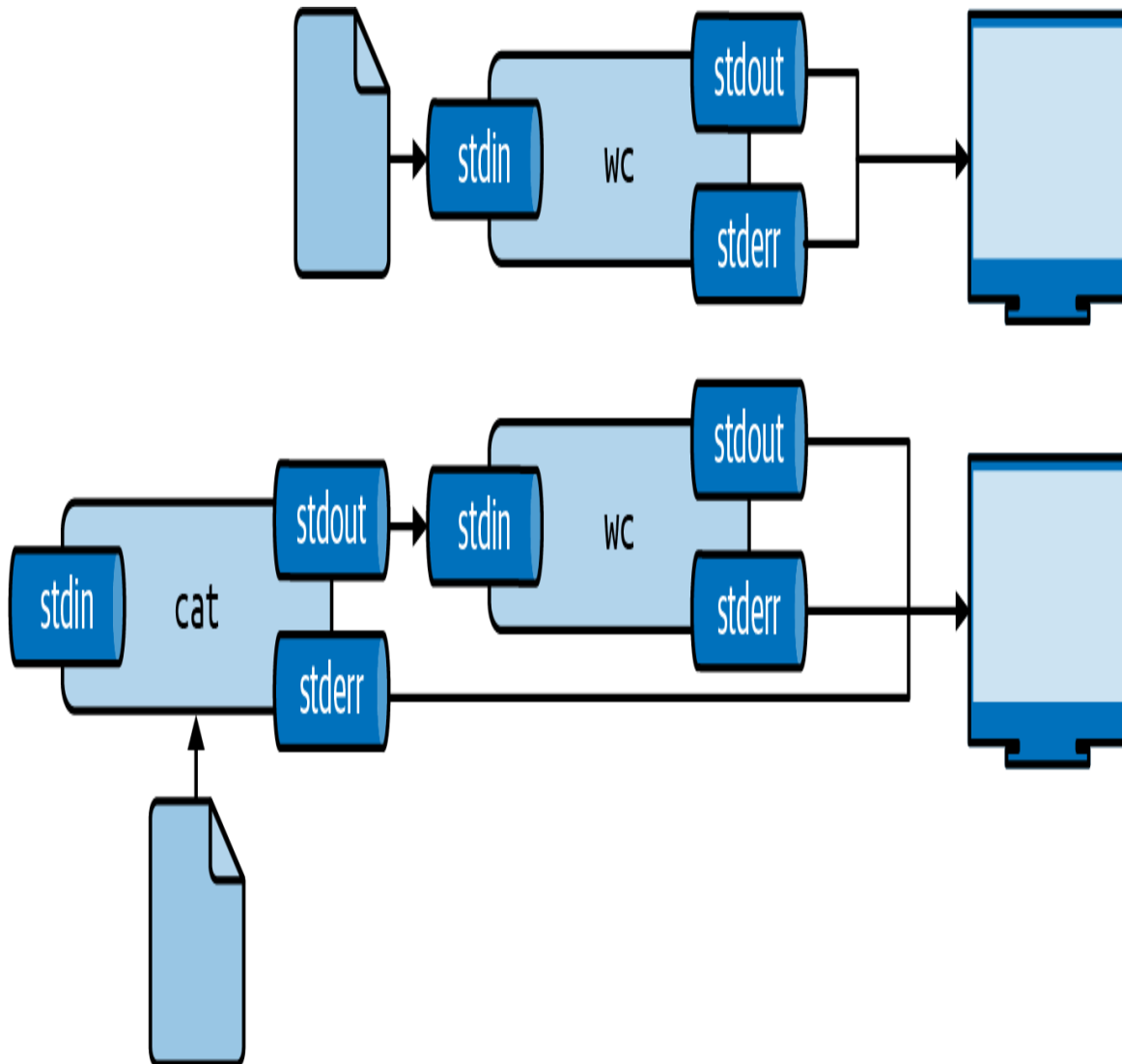


Figure 2-5. Two ways to use the contents of a file as input

Like many command-line tools, `wc` allows one or more filenames to be specified as arguments—for example:

```
$ wc -w greeting.txt movies.txt
2 greeting.txt
11 movies.txt
13 total
```

Note that in this case, `wc` also outputs the names of the files.

You can suppress the output of any tool by redirecting it to a special file called `/dev/null`. I often do this to suppress error messages (see [Figure 2-6](#) for an illustration). The following causes `cat` to produce an error message because it cannot find the file `404.txt`:

```
$ cat movies.txt 404.txt
Matrix
Star Wars
Home Alone
Indiana Jones
Back to the Future
cat: 404.txt: No such file or directory
```

You can redirect standard error to `/dev/null` as follows:

```
$ cat movies.txt 404.txt 2> /dev/null ❶
Matrix
Star Wars
Home Alone
Indiana Jones
Back to the Future
```

- ❶ The 2 refers to standard error.

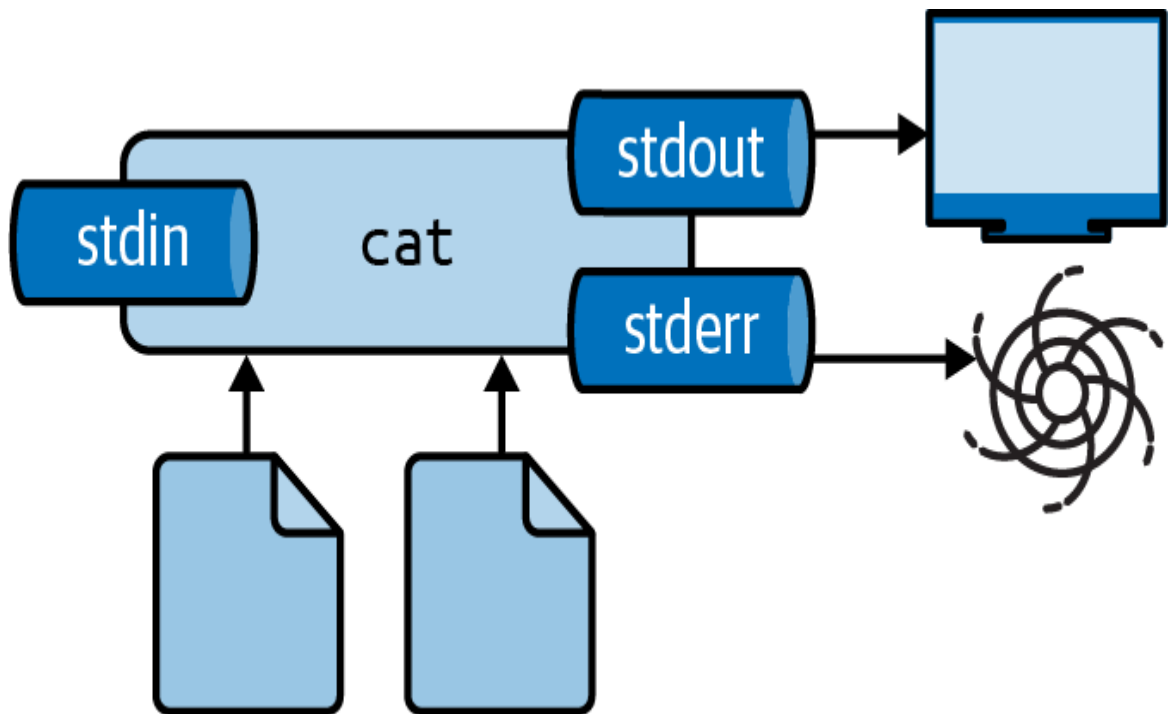


Figure 2-6. Redirecting `stderr` to `/dev/null`

Be careful not to read from and write to the same file. If you do, you'll end up with an empty file. That's because the tool whose output is redirected immediately opens that file for writing, thereby emptying it. There are two work-arounds for this: (1) write to a different file and rename it afterward with `mv`, or (2) use `sponge`,¹⁵ which soaks up all its input before writing to a file. [Figure 2-7](#) illustrates how this works.

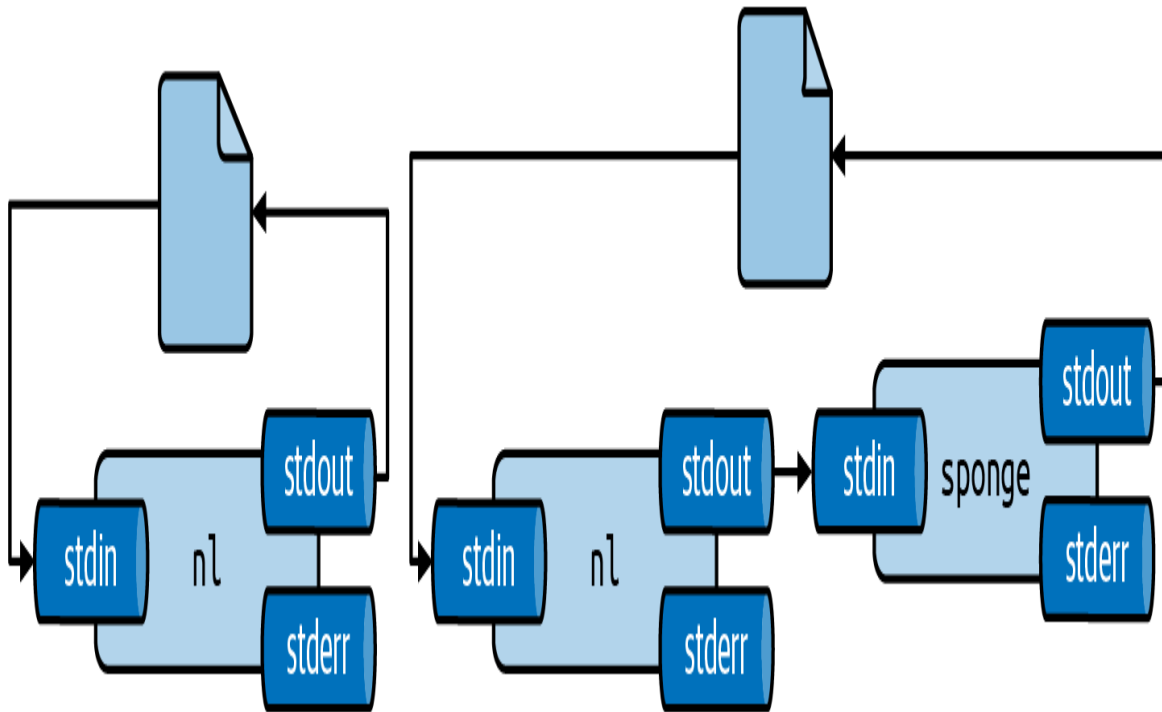


Figure 2-7. Unless you use `sponge`, you cannot read from and write to the same file in one pipeline

For example, imagine that you have used `dseq`¹⁶ to generate a file `dates.txt`, and now you'd like to add line numbers using `nl`.¹⁷ If you run the following, the file `dates.txt` will end up empty:

```
$ dseq 5 > dates.txt
$ < dates.txt nl > dates.txt
$ bat dates.txt
-----
| File: dates.txt  <EMPTY>
-----
```

Instead, you can use one of the work-arounds I just described:

```
$ dseq 5 > dates.txt
$ < dates.txt nl > dates-nl.txt
$ bat dates-nl.txt
-----
```

```
—
|   | File: dates-nl.txt
—
—
|   1  2021-06-30
|   2  2021-07-01
|   3  2021-07-02
|   4  2021-07-03
|   5  2021-07-04
—
—
$ dseq 5 > dates.txt
$ < dates.txt nl | sponge dates.txt
$ bat dates.txt
—
|
—
|   | File: dates.txt
—
—
|   1  2021-06-30
|   2  2021-07-01
|   3  2021-07-02
|   4  2021-07-03
|   5  2021-07-04
—
—
```

Working with Files and Directories

As data scientists, we work with a lot of data. This data is often stored in files. It is important to know how to work with files (and the directories they live in) on the command line. Every action that you can do using a GUI can be done with command-line tools (and you can do much more than that). In this section I introduce the most important tools to list, create, move, copy, rename, and delete files and directories.

Listing the contents of a directory can be done with `ls`. If you don't specify a directory, it lists the contents of the current directory. I prefer `ls` to have a long listing format and to have the directories grouped before files. Instead of typing the corresponding options each time, I use the alias `l`:

```
$ ls /data/ch10
alice.txt  count.py  count.R  Untitled1337.ipynb

$ alias l
l='ls --color -lhF --group-directories-first'

$ l /data/ch10
total 176K
-rw-r--r-- 1 dst dst 164K Jun 29 14:25 alice.txt
-rwxr-xr-x 1 dst dst  408 Jun 29 14:25 count.py*
-rw-r--r-- 1 dst dst  460 Jun 29 14:25 count.R
-rw-r--r-- 1 dst dst 1.7K Jun 29 14:25 Untitled1337.ipynb
```

You have already seen how we can create new files by redirecting the output with either `>` or `>>`. If you need to move a file to a different directory, you can use `mv`:¹⁸

```
$ mv hello.txt /data/ch02
```

You can also rename files with `mv`:

```
$ cd data
$ mv hello.txt bye.txt
```

You can also rename or move entire directories. If you no longer need a file, you can delete (or remove) it with `rm`:¹⁹

```
$ rm bye.txt
```

If you want to remove an entire directory with all its contents, specify the `-r` option, which stands for “recursive”:

```
$ rm -r /data/ch02/old
```

If you want to copy a file, use `cp`.²⁰ This is useful for creating backups:

```
$ cp server.log server.log.bak
```

You can create directories using `mkdir`.²¹

```
$ cd /data  
$ mkdir logs  
  
$ l  
total 44K  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch01/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch02/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch03/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch04/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch05/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch06/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch07/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch08/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch09/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 ch10/  
drwxr-xr-x 2 dst dst 4.0K Jun 29 14:25 logs/
```

TIP

Using the command-line tools to manage your files can be scary at first, because you have no graphical overview of the filesystem to provide immediate feedback. There are a few visual file managers that can help with this, such as GNU Midnight Commander, Ranger, and Vifm. These are not installed in the Docker image, but you can install one of them yourself by running `sudo apt install` followed by either `mc`, `ranger`, or `vifm`.

All of these command-line tools accept the `-v` option, which stands for *verbose*, so that they output what's going on. For example:

```
$ mkdir -v backup  
mkdir: created directory 'backup'
```

All tools other than `mkdir` also accept the `-i` option, which stands for *interactive*, and which causes the tools to ask you for confirmation. For example:

```
$ rm -i *  
zsh: sure you want to delete all 12 files in /data [yn]? n
```

Managing Output

Sometimes a tool or sequence of tools produces too much output to include in the book. Instead of manually altering such output, I prefer to be transparent by piping it through a helper tool. You don't necessarily have to do this, especially if you're interested in the complete output.

Here are the tools I use to make output manageable.

I often use `trim` to limit the output to a given height and width. By default, output is trimmed to 10 lines and the width of the terminal. Pass a negative number to disable trimming the height and/or the width. For example:

```
$ cat /data/ch07/tips.csv | trim 5 25
bill,tip,sex,smoker,day,....
16.99,1.01,Female,No,Sun...
10.34,1.66,Male,No,Sun,D...
21.01,3.5,Male,No,Sun,Di...
23.68,3.31,Male,No,Sun,D...
... with 240 more lines
```

Other tools that I use to massage the output are `head`, `tail`, `fold`, `paste`, and `column`. The Appendix contains an example for each of these.

If a file or an output contains a comma-separated value, I often pipe it through `csvlook` to turn it into a nice-looking table. If you run `csvlook`, you'll see the complete table. I have redefined `csvlook` such that the table is shortened by `trim`:

```
$ which csvlook
csvlook () {
    /usr/bin/csvlook "$@" | trim | sed 's/-| -/---/g;s/| -/|---/g;s/-|/
-|;s/|||/g;2s/-/-/g'
}

$ csvlook /data/ch07/tips.csv
+-----+-----+-----+-----+-----+-----+-----+
| bill | tip | sex | smoker | day | time | size |
+-----+-----+-----+-----+-----+-----+-----+
| 16.99 | 1.01 | Female | False | Sun | Dinner | 2 |
| 10.34 | 1.66 | Male | False | Sun | Dinner | 3 |
| 21.01 | 3.50 | Male | False | Sun | Dinner | 3 |
```

23.68	3.31	Male	False	Sun	Dinner	2
24.59	3.61	Female	False	Sun	Dinner	4
25.29	4.71	Male	False	Sun	Dinner	4
8.77	2.00	Male	False	Sun	Dinner	2
26.88	3.12	Male	False	Sun	Dinner	4

... with 236 more lines

I use `bat` to show the contents of a file where line numbers and syntax highlighting matter—for example:

```
$ bat /data/ch04/stream.py
-----
| File: /data/ch04/stream.py
-----
|
1 | #!/usr/bin/env python
2 | from sys import stdin, stdout
3 | while True:
4 |     line = stdin.readline()
5 |     if not line:
6 |         break
7 |     stdout.write("%d\n" % int(line)**2)
8 |     stdout.flush()
```

—

Sometimes I add the `-A` option when I want to explicitly point out the spaces, tabs, and newlines in a file.

Occasionally it's useful to write intermediate output to a file. This allows you to inspect any step in your pipeline once it has completed. You can insert the tool `tee` as often as you like in your pipeline. I often use it to inspect a portion of the final output, while writing the complete output to file (see [Figure 2-8](#)). Here, the complete output is written to `even.txt`, and the first five lines are printed using `trim`:

```
$ seq 0 2 100 | tee even.txt | trim 5
0
2
4
6
```

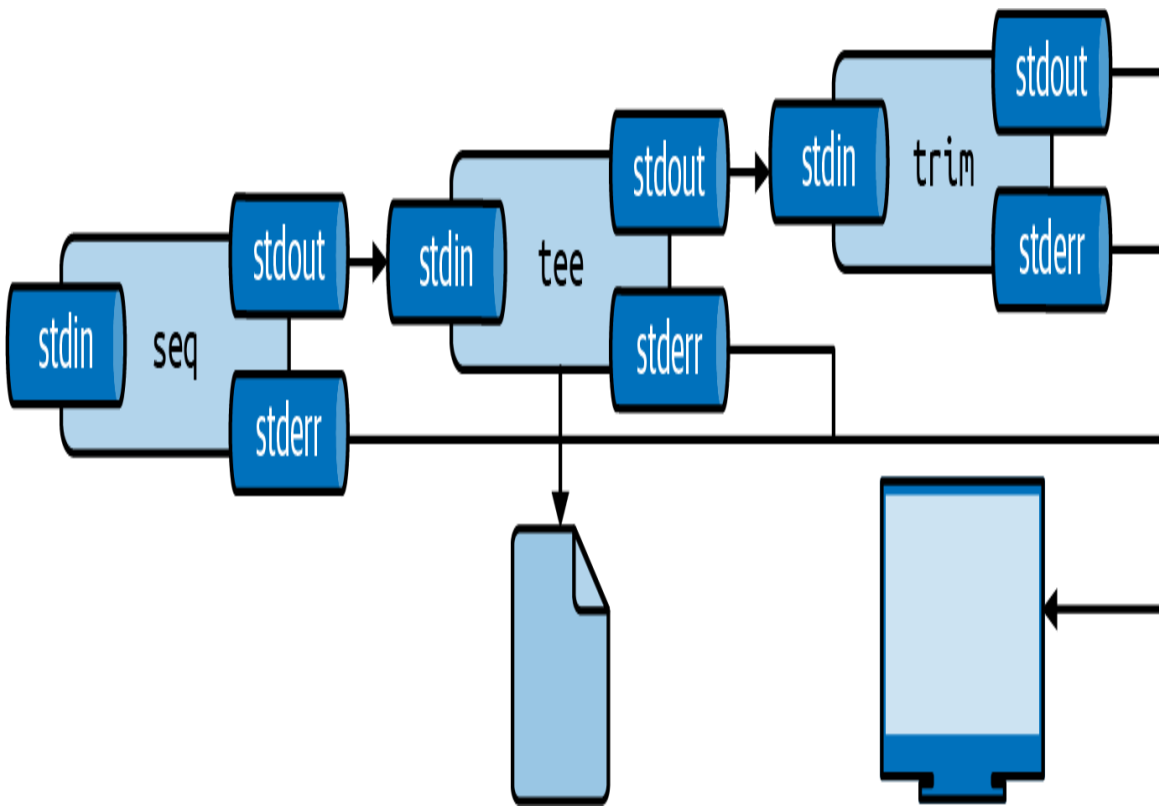


Figure 2-8. With `tee`, you can write intermediate output to a file

Last, to insert images that have been generated by command-line tools (that is, every image other than screenshots and diagrams) I use `display`. If you run `display`, you'll find that it doesn't work. In [Chapter 7](#), I explain four options for displaying images generated from the command line.

Help!

As you're finding your way around the command line, it may happen that you need help. Even the most seasoned users need help at some point. It is impossible to remember all the different command-line tools and their possible arguments. Fortunately, the command line offers several ways to get help.

Perhaps the most important command for getting help is `man`,²² which is short for *manual*. It contains information for most command-line tools. In

case I've forgotten the options to the tool `tar`, which happens all the time, I just access its manual page using the following:

```
$ man tar | trim 20
TAR(1)                               GNU TAR Manual                               TAR(1)

NAME
    tar - an archiving utility

SYNOPSIS
    Traditional usage
        tar {A|c|d|r|t|u|x}[GnSkUW0mpsMBiajJzZhPlRvwo] [ARG...]

    UNIX-style usage
        tar -A [OPTIONS] ARCHIVE ARCHIVE

        tar -c [-f ARCHIVE] [OPTIONS] [FILE...]
        tar -d [-f ARCHIVE] [OPTIONS] [FILE...]
        tar -t [-f ARCHIVE] [OPTIONS] [MEMBER...]
        tar -r [-f ARCHIVE] [OPTIONS] [FILE...]

... with 1147 more lines
```

Not every command-line tool has a manual page. Take `cd`, for example:

```
$ man cd
No manual entry for cd
```

For shell builtins like `cd`, you can consult the `zshbuiltins` manual page:

```
$ man zshbuiltins | trim
ZSHBUILTINS(1)                         General Commands Manual                  ZSHBUILTINS(1)

NAME
    zshbuiltins - zsh built-in commands

SHELL BUILTIN COMMANDS
    Some shell builtin commands take options as described in individual entries; these are often referred to in the list below as 'flags' to avoid confusion with shell options, which may also have an effect on
```

```
the behavior of builtin commands. In this introductory section, `op-
... with 2735 more lines
```

You can search by pressing / and exit by pressing q. Try to find the appropriate section for `cd`.

Newer command-line tools often lack a manual page as well. In such cases, your best bet is to invoke the tool with the `--help` (or `-h`) option. For example:

```
$ jq --help | trim
jq - commandline JSON processor [version 1.6]

Usage: jq [options] <jq filter> [file...]
      jq [options] --args <jq filter> [strings...]
      jq [options] --jsonargs <jq filter> [JSON_TEXTS...]

jq is a tool for processing JSON inputs, applying the given filter to
its JSON text inputs and producing the filter's results as JSON on
standard output.

... with 37 more lines
```

Specifying the `--help` option also works for command-line tools such as `cat`. However, the corresponding manual page often provides more information. If, after trying these three approaches, you are still stuck, then consulting the internet is perfectly acceptable. In the [Appendix](#), there's a list of all the command-line tools used in this book. Besides showing how each command-line tool can be installed, the Appendix also shows how you can get help for each tool.

Manual pages can be quite verbose and difficult to read. The tool `tldr`²³ (which is short for “too long; didn't read”) is a collection of community-maintained help pages for command-line tools that aims to be a simpler, more approachable complement to traditional manual pages. Here's an example of the `tldr` page for `tar`:

```
$ tldr tar | trim 20

tar
```

Archiving utility.

Often combined with a compression method, such as gzip or bzip2.

More information: <https://www.gnu.org/software/tar/>.

- [c]reate an archive and write it to a [f]ile:
`tar cf target.tar file1 file2 file3`
- [c]reate a g[z]ipped archive and write it to a [f]ile:
`tar czf target.tar.gz file1 file2 file3`
- [c]reate a g[z]ipped archive from a directory using relative paths:
`tar czf target.tar.gz --directory=path/to/directory .`
- E[x]tract a (compressed) archive [f]ile into the current directory
[v]erbos...
`tar xvf source.tar[.gz|.bz2|.xz]`
 - E[x]tract a (compressed) archive [f]ile into the target directory:
... with 12 more lines

As you can see, rather than listing the many options alphabetically like `man` often does, `tldr` cuts to the chase by giving you a list of practical examples.

Summary

In this chapter you learned how to get all the required command-line tools by installing a Docker image. I also went over some essential command-line concepts and how to get help. Now that you have all the necessary ingredients, you're ready for the first step of the OSEMN model for data science: obtaining data.

For Further Exploration

- The subtitle of this book pays homage to the epic *Unix Power Tools, 3rd ed.* by Shelley Powers, Jerry Peek, Tim O'Reilly, and Mike Loukides (O'Reilly), and rightly so. With over 51 chapters and more than a thousand pages, *Unix Power Tools* covers just

about everything there is to know about Unix. It weighs more than four pounds, so you might want to consider getting the ebook.

- The website [explainshell](#) parses a command or a sequence of commands and provides a short explanation of each part. This site is useful for quickly understanding a new command or option without having to skim through the relevant manual pages.
- Docker is truly a brilliant piece of software. In this chapter I've briefly explained how to download a Docker image and run a Docker container, but it might be worthwhile to [learn how to create your own Docker images](#). The book *Docker: Up & Running*, 2nd ed. by Sean P. Kane and Karl Matthias (O'Reilly) is a good resource as well.

1 Richard M. Stallman and David MacKenzie, *ls – List Directory Contents*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

2 Torbjorn Granlund and Richard M. Stallman, *cat – Concatenate Files and Print on the Standard Output*, version 8.30, 2018, <https://www.gnu.org/software/coreutils>.

3 Stephen Dolan, *jq – Command-Line JSON Processor*, version 1.6, 2021, <https://stedolan.github.io/jq/>

4 Ulrich Drepper, *seq – Print a Sequence of Numbers*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

5 Jim Meyering, *pwd – Print Name of Current/Working Directory*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

6 David MacKenzie and Jim Meyering, *head – Output the First Part of Files*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

7 David M. Ihnat and David MacKenzie, *paste – Merge Lines of Files*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

8 Philip A. Nelson, *bc – an Arbitrary Precision Calculator Language*, version 1.07.1, 2017, <https://www.gnu.org/software/bc>.

9 Eric S. Raymond, *The Art of Unix Programming* (Addison-Wesley).

10 Jim Meyering, *grep – Print Lines That Match Patterns*, version 3.4, 2019, <https://www.gnu.org/software/grep>.

11 Paul Rubin and David MacKenzie, *wc – Print Newline, Word, and Byte Counts for Each File*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

- 12 Mike Haertel and Paul Eggert, *sort – Sort Lines of Text Files*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 13 Karel Zak, *rev – Reverse Lines Characterwise*, version 2.36.1, 2021, <https://www.kernel.org/pub/linux/utils/util-linux>.
- 14 Some consider this a “useless use” of `cat`, arguing that the purpose of `cat` is to concatenate files, and not using it for that purpose is a waste of time and costs you a process. I think this is silly. We’ve got more important things to do!
- 15 Colin Watson and Tollef Fog Heen, *sponge – Soak Up Standard Input and Write to a File*, version 0.65, 2021, <https://joeyh.name/code/moreutils>.
- 16 Jeroen Janssens, *dseq – Generate Sequence of Dates*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.
- 17 Scott Bartram and David MacKenzie, *nl – Number Lines of Files*, version 8.30, 2020, <https://www.gnu.org/software/coreutils>.
- 18 Mike Parker, David MacKenzie, and Jim Meyering, *mv – Move (Rename) Files*, version 8.30, 2020, <https://www.gnu.org/software/coreutils>.
- 19 Paul Rubin et al., *rm – Remove Files or Directories*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 20 Torbjorn Granlund, David MacKenzie, and Jim Meyering, *cp – Copy Files and Directories*, version 8.30, 2018, <https://www.gnu.org/software/coreutils>.
- 21 David MacKenzie, *mkdir – Make Directories*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 22 John W. Eaton and Colin Watson, *man – an Interface to the System Reference Manuals*, version 2.9.1, 2020, <https://nongnu.org/man-db>.
- 23 Owen Voke, *tldr – Collaborative Cheatsheets for Console Commands*, version 3.3.7, 2021, <https://tldr.sh>.

Chapter 3. Obtaining Data

This chapter deals with the first step of the OSEMN model: obtaining data. After all, without any data, there is not much data science that we can do. I assume that the data you need to solve your data science problem already exists. Your first task is to get this data onto your computer (and possibly also inside the Docker container) in a form that you can work with.

According to the Unix philosophy, text is a universal interface. Almost every command-line tool takes text as input, produces text as output, or both. This is the main reason why command-line tools can work so well together. However, as we'll see, even just text can come in multiple forms.

Data can be obtained in several ways—for example, by downloading it from a server, querying a database, or connecting to a Web API. Sometimes the data comes in a compressed form or in a binary format such as a Microsoft Excel Spreadsheet. In this chapter, I discuss several tools that help tackle this from the command line, including `curl`,¹ `in2csv`,² `sql2csv`,³ and `tar`.⁴

Overview

In this chapter, you'll learn how to:

- Copy local files to the Docker image
- Download data from the internet
- Decompress files
- Extract data from spreadsheets
- Query relational databases
- Call web APIs

This chapter starts with the following files:

```
$ cd /data/ch03  
$ l  
total 924K  
-rw-r--r-- 1 dst dst 627K Jun 29 14:26 logs.tar.gz  
-rw-r--r-- 1 dst dst 189K Jun 29 14:26 r-datasets.db  
-rw-r--r-- 1 dst dst 149 Jun 29 14:26 tmnt-basic.csv  
-rw-r--r-- 1 dst dst 148 Jun 29 14:26 tmnt-missing-newline.csv  
-rw-r--r-- 1 dst dst 181 Jun 29 14:26 tmnt-with-header.csv  
-rw-r--r-- 1 dst dst 91K Jun 29 14:26 top2000.xlsx
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Copying Local Files to the Docker Container

A common situation is that you already have the necessary files on your own computer. This section explains how you can get those files into the Docker container.

I mentioned in [Chapter 2](#) that the Docker container is an isolated virtual environment. Luckily, there is one exception to that: files can be transferred in and out of the Docker container. The local directory from which you ran `docker run` is mapped to a directory in the Docker container. This directory is called `/data`. Note that this is not the home directory, which is `/home/dst`.

If you have one or more files on your local computer, and you want to apply some command-line tools to them, all you have to do is copy or move the files to that mapped directory. Let's assume that you have a file called `logs.csv` in your Downloads directory.

If you're running Windows, open the Command Prompt or PowerShell and run the following two commands:

```
> cd %UserProfile%\Downloads  
> copy logs.csv MyDataScienceToolbox\
```

If you are running Linux or macOS, open a terminal and execute the following command on your operating system (not inside the Docker container):

```
$ cp ~/Downloads/logs.csv ~/my-data-science-toolbox
```

You can also drag and drop the file into the right directory using a graphical file manager such as Windows Explorer or macOS Finder.

Downloading from the Internet

The internet provides, without a doubt, the largest resource for interesting data. The command-line tool `curl` can be considered the command line's Swiss Army knife when it comes to downloading data from the internet.

Introducing curl

When you browse a URL, which stands for *uniform resource locator*, your browser interprets the data it downloads. For example, the browser renders HTML files, plays video files automatically, and shows PDF files.

However, when you use `curl` to access a URL, it downloads the data and, by default, prints it to standard output. `curl` doesn't do any interpretation, but luckily other command-line tools can be used to process the data further.

The easiest invocation of `curl` is to specify a URL as a command-line argument. Let's try downloading an article from Wikipedia:

```
$ curl "https://en.wikipedia.org/wiki/List_of_windmills_in_the_Netherlands" |  
> trim ①  
% Total    % Received % Xferd  Average Speed   Time     Time     Time  
Current                                         Dload  Upload   Total   Spent   Left   Speed  
0       0      0      0      0      0      0      0 --::-- --::-- --::--  
0<!DOCTYPE html>  
<html class="client-nojs" lang="en" dir="ltr">  
<head>
```

```
<meta charset="UTF-8"/>
<title>List of windmills in the Netherlands - Wikipedia</title>
<script>document.documentElement.className="client-js";RLCONF=
{"wgBreakFrames":...
"wikitext","wgRelevantPageName":"List_of_windmills_in_the_Netherlands","wgRele
v...
"site.styles":"ready","noscript":"ready","user.styles":"ready","ext.globalCssJ
s...
"ext.growthExperiments.SuggestedEditSession"];</script>
<script>(RLQ=window.RLQ||[]).push(function()
{mw.loader.implement("user.options@...
100 244k 100 244k 0 0 1291k 0 --:---- --:--- --:--- 1291k
... with 1723 more lines
```

- ❶ Remember, `trim` is used only to make the output fit nicely in the book.

As you can see, `curl` downloads the raw HTML returned by Wikipedia's server; no interpretation is being done, and the contents are immediately printed on standard output. Because of the URL, you'd think that this article would list all the windmills in the Netherlands. However, there are apparently so many windmills that each province has its own page. Fascinating.

By default, `curl` outputs a progress meter that shows the download rate and the expected time of completion. This output isn't written to standard output but instead is written to a separate channel known as *standard error*, so that it doesn't interfere when you add another tool to the pipeline. While this information can be useful when downloading very large files, I usually find it distracting, so I specify the `-s` option to *silence* this output:

```
$ curl -s "https://en.wikipedia.org/wiki/List_of_windmills_in_Friesland" |
> pup -n 'table.wikititable tr' ❶
234
```

- ❶ I'll discuss `pup`,⁵ a handy tool for scraping websites, in more detail in [Chapter 5](#).

And what do you know, there are apparently 234 windmills in the province of Friesland alone!

Saving

You can let `curl` save the output to a file by adding the `-O` option. The filename will be based on the last part of the URL:

```
$ curl -s "https://en.wikipedia.org/wiki/List_of_windmills_in_Friesland" -O  
  
$ ls  
total 1.4M  
-rw-r--r-- 1 dst dst 427K Jun 29 14:27 List_of_windmills_in_Friesland  
-rw-r--r-- 1 dst dst 627K Jun 29 14:26 logs.tar.gz  
-rw-r--r-- 1 dst dst 189K Jun 29 14:26 r-datasets.db  
-rw-r--r-- 1 dst dst 149 Jun 29 14:26 tmnt-basic.csv  
-rw-r--r-- 1 dst dst 148 Jun 29 14:26 tmnt-missing-newline.csv  
-rw-r--r-- 1 dst dst 181 Jun 29 14:26 tmnt-with-header.csv  
-rw-r--r-- 1 dst dst 91K Jun 29 14:26 top2000.xlsx
```

If you don't like that filename, then you can use the `-o` option together with a filename or redirect the output to a file yourself:

```
$ curl -s "https://en.wikipedia.org/wiki/List_of_windmills_in_Friesland" >  
fries  
land.html
```

Other Protocols

In total, `curl` supports **more than 20 protocols**. To download from an FTP server (FTP stands for “File Transfer Protocol”), you use `curl` the same way. Here I download the file *welcome.msg* from *ftp.gnu.org*:

```
$ curl -s "ftp://ftp.gnu.org/welcome.msg" | trim  
NOTICE (Updated December 18 2018):  
  
FSF public IP addresses are changing between December 20 and January 7th  
  
If you have hardcoded the IP address of any GNU/FSF servers in those  
ranges in any code or configuration files, they will need to be  
updated. If you refer to our servers by their DNS name, such as
```

```
"gnu.org", then that will continue to work. You should use the DNS name wherever possible.
```

```
... with 68 more lines
```

If the specified URL is a directory, `curl` will list the contents of that directory. When the URL is password protected, you can specify a username and a password with the `-u` option.

Or there's the DICT protocol, which allows you to access various dictionaries and look up definitions. Here's the definition of *windmill* according to the Collaborative International Dictionary of English:

```
$ curl -s "dict://dict.org/d:windmill" | trim
220 dict.dict.org dictd 1.12.1/rf on Linux 4.19.0-10-amd64 <auth.mime>
<4623255...
250 ok
150 1 definitions retrieved
151 "Windmill" gcide "The Collaborative International Dictionary of English
v.0...
Windmill \Wind"mill`\, n.
A mill operated by the power of the wind, usually by the
action of the wind upon oblique vanes or sails which radiate
from a horizontal shaft. --Chaucer.
[1913 Webster]
.
... with 2 more lines
```

When you are downloading data from the internet, however, the protocol will most likely be HTTP, so the URL will start with either `http://` or `https://`.

Following Redirects

When you access a shortened URL, such as the one that starts with `http://bit.ly/` or `http://t.co/`, your browser automatically redirects you to the correct location. With `curl`, however, you need to specify the `-L` or `--location` option in order to be redirected. If you don't, you can get something like:

```
$ curl -s "https://bit.ly/2XBxvwK"
<html>
<head><title>Bitly</title></head>
<body><a href="https://youtu.be/dQw4w9WgXcQ">moved here</a></body>
</html>%
```

Sometimes you get nothing back, like when we follow the URL just mentioned:

```
$ curl -s "https://youtu.be/dQw4w9WgXcQ"
```

If you specify the `-I` or `--head` option, `curl` fetches only the HTTP header of the response, which allows you to inspect the status code and other information returned by the server:

```
$ curl -sI "https://youtu.be/dQw4w9WgXcQ" | trim
HTTP/2 303
content-type: application/binary
x-content-type-options: nosniff
cache-control: no-cache, no-store, max-age=0, must-revalidate
pragma: no-cache
expires: Mon, 01 Jan 1990 00:00:00 GMT
date: Tue, 29 Jun 2021 12:27:13 GMT
location: https://www.youtube.com/watch?v=dQw4w9WgXcQ&feature=youtu.be
content-length: 0
x-frame-options: SAMEORIGIN
... with 9 more lines
```

The first line shows the protocol followed by the HTTP status code, which is 303 in this case. You can also see the location this URL redirects to.

Inspecting the header and getting the status code is a useful debugging tool in case `curl` does not give you the expected result. Other common HTTP status codes include 404 (not found) and 403 (forbidden). Wikipedia has a page that lists [all HTTP status codes](#).

In summary, `curl` is a useful command-line tool for downloading data from the internet. Its three most common options are `-s` to silence the progress meter, `-u` to specify a username and password, and `-L` to automatically follow redirects. See the man page for more information (and to make your head spin):

```
$ man curl | trim 20
curl(1)                               Curl Manual                  curl(1)

NAME
    curl - transfer a URL

SYNOPSIS
    curl [options / URLs]

DESCRIPTION
    curl is a tool to transfer data from or to a server, using one of the
    supported protocols (DICT, FILE, FTP, FTPS, GOPHER, HTTP, HTTPS, IMAP,
    IMAPS, LDAP, LDAPS, MQTT, POP3, POP3S, RTMP, RTMPS, RTSP, SCP, SFTP,
    SMB, SMBS, SMTP, SMTPS, TELNET and TFTP). The command is designed to
    work without user interaction.

    curl offers a busload of useful tricks like proxy support, user authentication,
    FTP upload, HTTP post, SSL connections, cookies, file transfer resume,
    Metalink, and more. As you will see below, the number of features will make your head spin!

... with 3986 more lines
```

Decompressing Files

If the original dataset is very large or is a collection of many files, it may be a compressed archive. Datasets that contain many repeated values (such as the words in a text file or the keys in a JSON file) are especially well suited for compression.

Common file extensions of compressed archives are *.tar.gz*, *.zip*, and *.rar*. To decompress these, you would use the command-line tools `tar`, `unzip`,⁶ and `unrar`,⁷ respectively. (There are a few less-common file extensions for which you would need other tools.)

Let's take *tar.gz* (pronounced “gzipped tarball”) as an example. To extract an archive named *logs.tar.gz*, you would use the following incantation:

```
$ tar -xzf logs.tar.gz ❶
```

❶

It's common to combine these three short options, like I did here, but you can also specify them separately as `-x` `-z` `-f`. In fact, many command-line tools allow you to combine options that consist of a single character.

Indeed, `tar` is notorious for its many command-line arguments. In this case, the three options `-x`, `-z`, and `-f` specify that `tar` should *extract* files from an archive, use gzip as the decompression algorithm, and use the file `logs.tar.gz`.

However, since we're not yet familiar with this archive, it's a good idea to first examine its contents. This can be done using the `-t` option (instead of the `-x` option):

```
$ tar -tzf logs.tar.gz | trim  
E1F0SPSAYDNUZI.2020-09-01-00.0dd00628  
E1F0SPSAYDNUZI.2020-09-01-00.b717c457  
E1F0SPSAYDNUZI.2020-09-01-01.05f904a4  
E1F0SPSAYDNUZI.2020-09-01-02.36588daf  
E1F0SPSAYDNUZI.2020-09-01-02.6cea8b1d  
E1F0SPSAYDNUZI.2020-09-01-02.be4bc86d  
E1F0SPSAYDNUZI.2020-09-01-03.16f3fa32  
E1F0SPSAYDNUZI.2020-09-01-03.1c0a370f  
E1F0SPSAYDNUZI.2020-09-01-03.76df64bf  
E1F0SPSAYDNUZI.2020-09-01-04.0a1ade1b  
... with 2427 more lines
```

It seems that this archive contains a lot of files, and they are not inside a directory. To keep the current directory clean, it's a good idea to first create a new directory using `mkdir` and extract those files there using the `-C` option:

```
$ mkdir logs  
$ tar -xzf logs.tar.gz -C logs
```

Let's verify the number of files and some of their contents:

```
$ ls logs | wc -l
2437

$ cat logs/* | trim
#Version: 1.0
#Fields: date time x-edge-location sc-bytes c-ip cs-method cs(Host) cs-uri-
stem...
2020-09-01      00:51:54          SEA19-C1        391    206.55.174.150  GET
...
2020-09-01      00:54:59          CPH50-C2        384    82.211.213.95  GET
...
#Version: 1.0
#Fields: date time x-edge-location sc-bytes c-ip cs-method cs(Host) cs-uri-
stem...
2020-09-01      00:04:28          DFW50-C1        391
2a03:2880:11ff:9::face:...
#Version: 1.0
#Fields: date time x-edge-location sc-bytes c-ip cs-method cs(Host) cs-uri-
stem...
2020-09-01      01:04:14          ATL56-C4        385
2600:1700:2760:da20:548...
... with 10279 more lines
```

Excellent. Now, I understand that you'd like to scrub and explore these log files, but we'll get to that later, in [Chapter 5](#) and [Chapter 7](#).

In time you'll get used to these options, but I'd like to show you an alternative that you might find convenient. Rather than you having to remember the different command-line tools and their options, there's a handy script called `unpack`⁸ that will decompress many different formats. `unpack` looks at the extension of the file that you want to decompress and calls the appropriate command-line tool. Now, in order to decompress this same file, you would run:

```
$ unpack logs.tar.gz
```

Converting Microsoft Excel Spreadsheets to CSV

For many people, Microsoft Excel offers an intuitive way to work with small datasets and perform calculations on them. As a result, a lot of data is embedded into Microsoft Excel spreadsheets. These spreadsheets are, depending on the extension of the filename, stored either in a proprietary binary format (.xls) or as a collection of compressed XML files (.xlsx). In both cases, the data is not readily usable by most command-line tools. It would be a shame if we could not use those valuable datasets just because they are stored this way.

Especially when you’re just starting out at the command line, you might be tempted to convert your spreadsheet to CSV by opening it in Microsoft Excel, or in an open source variant such as LibreOffice Calc, and manually exporting it to CSV. While this works as a one-off solution, it does not scale well to multiple files and cannot be automated. Furthermore, when you’re working on a server, chances are that you don’t have such an application available. Trust me, you’ll get the hang of it.

Luckily, there is a command-line tool called `in2csv` that converts Microsoft Excel spreadsheets to CSV files (CSV stands for “comma-separated values”). Working with CSV can be tricky because it lacks a formal specification. Yakov Shafranovich defines the CSV format according to the following three points:⁹

1. Each record is located on a separate line, delimited by a line break (`\n`). Take, for example, the following CSV file with crucial information about the Teenage Mutant Ninja Turtles:

```
$ bat -A tmnt-basic.csv ①
-----
| File: tmnt-basic.csv
-----
1 | Leonardo,Leo,blue,two·ninjakens ↵
2 | Raphael,Raph,red,pair·of·sai ↵
```

```
3 | Michelangelo,Mikey.or.Mike,orange,pair.of.nunchaku ↵
4 | Donatello,Donnie.or.Don,purple,staff ↵
```

- ① The **-A** option makes **bat** show all nonprintable characters like spaces, tabs, and newlines.

2. The last record in the file may or may not have an ending line break. For example:

```
$ bat -A tmnt-missing-newline.csv
```

```
| File: tmnt-missing-newline.csv
|
1 | Leonardo,Leo,blue,two·nинжакенс ↵
2 | Raphael,Raph,red,pair.of.sai ↵
3 | Michelangelo,Mikey.or.Mike,orange,pair.of.nunchaku ↵
4 | Donatello,Donnie.or.Don,purple,staff
```

3. There may be a header appearing as the first line of the file with the same format as normal record lines. This header will contain names corresponding to the fields in the file and should contain the same number of fields as the records in the rest of the file. For example:

```
$ bat -A tmnt-with-header.csv
```

```
| File: tmnt-with-header.csv
|
1 | name,nickname,mask_color,weapon ↵
2 | Leonardo,Leo,blue,two·нинжакенс ↵
3 | Raphael,Raph,red,pair.of.sai ↵
4 | Michelangelo,Mikey.or.Mike,orange,pair.of.nunchaku ↵
5 | Donatello,Donnie.or.Don,purple,staff ↵
```

As you can see, CSV by default is not too readable. You can pipe the data to a tool called `csvlook`,¹⁰ which will nicely format it into a table. If the CSV data has no header, as in the file `tmnt-missing-newline.csv`, then you need to add the `-H` option; otherwise the first line will be interpreted as the header:

```
$ csvlook tmnt-with-header.csv
```

name	nickname	mask_color	weapon
Leonardo	Leo	blue	two ninjakens
Raphael	Raph	red	pair of sai
Michelangelo	Mikey or Mike	orange	pair of nunchaku
Donatello	Donnie or Don	purple	staff

```
$ csvlook tmnt-basic.csv
```

Leonardo	Leo	blue	two ninjakens
Raphael	Raph	red	pair of sai
Michelangelo	Mikey or Mike	orange	pair of nunchaku
Donatello	Donnie or Don	purple	staff

```
$ csvlook -H tmnt-missing-newline.csv
```

❶

a	b	c	d
Leonardo	Leo	blue	two ninjakens
Raphael	Raph	red	pair of sai
Michelangelo	Mikey or Mike	orange	pair of nunchaku
Donatello	Donnie or Don	purple	staff

- ❶ The `-H` option specifies that the CSV file has no header.

Let's demonstrate `in2csv` using a spreadsheet that contains the two thousand most popular songs according to an annual Dutch marathon radio program, the **Top 2000**. To extract the Excel file's data, you invoke `in2csv` as follows:

```
$ curl "https://www.nporadio2.nl/data/download/TOP-2000-2020.xlsx" > top2000.xls
```

X

```
% Total     % Received % Xferd  Average Speed   Time     Time     Time
Current                                         Dload  Upload   Total Spent  Left  Speed
100 92838  100 92838    0       0  625k      0 --::-- --::-- --::-- 625k
```

```
$ in2csv top2000.xlsx | tee top2000.csv | trim
NR.,ARTIEST,TITEL,JAAR
1,Danny Vera,Roller Coaster,2019
2,Queen,Bohemian Rhapsody,1975
3,Eagles,Hotel California,1977
4,Billy Joel,Piano Man,1974
5,Led Zeppelin,Stairway To Heaven,1971
6,Pearl Jam,Black,1992
7,Boudewijn de Groot,Avond,1997
8,Coldplay,Fix You,2005
9,Pink Floyd,Wish You Were Here,1975
... with 1991 more lines
```

Who is Danny Vera? The most popular song is supposed to be “Bohemian Rhapsody,” of course. Well, at least Queen appears plenty of times in the Top 2000, so I can’t really complain:

```
$ csvgrep top2000.csv --columns ARTIEST --regex '^Queen$' | csvlook -I ❶
+---+---+---+---+
| NR. | ARTIEST | TITEL | JAAR |
+---+---+---+---+
| 2 | Queen | Bohemian Rhapsody | 1975 |
| 11 | Queen | Love Of My Life | 1975 |
| 46 | Queen | Innuendo | 1991 |
| 55 | Queen | Don't Stop Me Now | 1979 |
| 70 | Queen | Somebody To Love | 1976 |
| 85 | Queen | Who Wants To Live Forever | 1986 |
| 89 | Queen | The Show Must Go On | 1991 |
| 131 | Queen | Killer Queen | 1974 |
... with 24 more lines
```

- ❶ The value after the `--regex` option is a regular expression (or *regex*). It’s a special syntax for defining patterns. Here, I only want to match artists that exactly match “Queen,” so I use the caret (^) and dollar sign (\$) to match the start and end of the values in the column *ARTIEST*.

By the way, the tools `in2csv`, `csvgrep`, and `csvlook` are part of `csvkit`, which is a collection of command-line tools for working with CSV data.

The format of the file is automatically determined by the extension (`.xlsx`, in this case). If you were to pipe the data into `in2csv`, you would have to specify the format explicitly.

A spreadsheet can contain multiple worksheets. `in2csv` extracts the first worksheet by default. To extract a different worksheet, you need to pass the name of the worksheet to the `--sheet` option. If you're not sure what the worksheet is called, you can use the `--names` option, which prints the names of all the worksheets. Here we see that `top2000.xlsx` has only one sheet, named *Blad1* (which is Dutch for Sheet1):

```
$ in2csv --names top2000.xlsx
Blad1
```

Querying Relational Databases

Many companies store their data in a relational database. Just as with spreadsheets, it would be great if we could obtain that data from the command line.

Examples of relational databases are MySQL, PostgreSQL, and SQLite. These databases all have a slightly different way of interfacing with them. Some provide a command-line tool or a command-line interface, while others do not. Moreover, they are not very consistent when it comes to their usage and output.

Fortunately, there is a command-line tool called `sql2csv` that is part of the `csvkit` suite and that works with many different databases through a common interface, including MySQL, Oracle, PostgreSQL, SQLite, Microsoft SQL Server, and Sybase. The output of `sql2csv` is, as its name suggests, in CSV format.

We can obtain data from relational databases by executing a `SELECT` query on them. (`sql2csv` also supports `INSERT`, `UPDATE`, and `DELETE` queries, but

that's not the focus of this chapter.)

`sql2csv` needs two arguments: `--db`, which specifies the database URL, of which the typical form is

`dialect+driver://username:password@host:port/database`; and `--query`, which contains the `SELECT` query. For example, given an SQLite database that contains the standard datasets from R,¹¹ I can select all the rows from the table `mtcars` and sort them by the `mpg` column as follows:

```
$ sql2csv --db 'sqlite:///r-datasets.db' \
> --query 'SELECT row_names AS car, mpg FROM mtcars ORDER BY mpg' | csvlook
| car           | mpg |
| Cadillac Fleetwood | 10.4 |
| Lincoln Continental | 10.4 |
| Camaro Z28       | 13.3 |
| Duster 360      | 14.3 |
| Chrysler Imperial | 14.7 |
| Maserati Bora   | 15.0 |
| Merc 450SLC     | 15.2 |
| AMC Javelin     | 15.2 |
... with 24 more lines
```

This SQLite database is a local file, so in this I don't need to specify any username, password, or host. If you wanted to query the database of your employer, you'd of course need to know how to access it, and you'd need permission to do so.

Calling Web APIs

In the previous section I explained how to download files from the internet. Another way data can come from the internet is through a web API. The number of APIs being offered is growing at an increasing rate, which means a lot of interesting data for us data scientists.

Web APIs are not meant to be presented in a nice layout, like a website. Instead, most web APIs return data in a structured format, such as JSON or XML. Having data in a structured form has the advantage that the data can be easily processed by other tools, such as `jq`. For example, an API of Ice

and Fire, which contains a lot of information about George R. R. Martin's fictional world (in which the *Game of Thrones* books and TV show take place) returns data in the following JSON structure:

```
$ curl -s "https://anapioficeandfire.com/api/characters/583" | jq '.'
{
  "url": "https://anapioficeandfire.com/api/characters/583",
  "name": "Jon Snow",
  "gender": "Male",
  "culture": "Northmen",
  "born": "In 283 AC",
  "died": "", ①
  "titles": [
    "Lord Commander of the Night's Watch"
  ],
  "aliases": [
    "Lord Snow",
    "Ned Stark's Bastard",
    "The Snow of Winterfell",
    "The Crow-Come-Over",
    "The 998th Lord Commander of the Night's Watch",
    "The Bastard of Winterfell",
    "The Black Bastard of the Wall",
    "Lord Crow"
  ],
  "father": "",
  "mother": "",
  "spouse": "",
  "allegiances": [
    "https://anapioficeandfire.com/api/houses/362"
  ],
  "books": [
    "https://anapioficeandfire.com/api/books/5"
  ],
  "povBooks": [
    "https://anapioficeandfire.com/api/books/1",
    "https://anapioficeandfire.com/api/books/2",
    "https://anapioficeandfire.com/api/books/3",
    "https://anapioficeandfire.com/api/books/8"
  ],
  "tvSeries": [
    "Season 1",
    "Season 2",
    "Season 3",
    "Season 4",
    "Season 5",
    "Season 6",
    "Season 7",
    "Season 8"
  ]
}
```

```
        "Season 6"  
    ],  
    "playedBy": [  
        "Kit Harington"  
    ]  
}
```

- ❶ Spoiler alert: this data is not entirely up to date.

The data is piped to the command-line tool `jq` just to display it in a nice way. `jq` has many more scrubbing and exploring possibilities that I will explore in [Chapter 5](#) and [Chapter 7](#).

Authentication

Some web APIs require you to authenticate (that is, to prove your identity) before you can consume their output. There are several ways to do this. Some web APIs use API keys, while others use the OAuth protocol. News API, an independent source of headlines and news articles, is a great example. Let's see what happens when you try to access this API without an API key:

```
$ curl -s "http://newsapi.org/v2/everything?q=linux" | jq -  
{  
    "status": "error",  
    "code": "apiKeyMissing",  
    "message": "Your API key is missing. Append this to the URL with the apiKey  
pa  
ram, or use the x-api-key HTTP header."  
}
```

Well, that was to be expected. The part after the question mark, by the way, is where we pass any query parameters. That's also the place where you need to specify an API key. I'd like to keep my own API key a secret, so I insert it below by reading the file `/data/.secret/newsapi.org_apikey` using command substitution:

```
$ curl -s "http://newsapi.org/v2/everything?q=linux&apiKey=$(  
/data/.secret/new  
sapi.org_apikey)" |  
> jq '.' | trim 30  
{  
    "status": "ok",  
    "totalResults": 9616,  
    "articles": [  
        {  
            "source": {  
                "id": "techcrunch",  
                "name": "TechCrunch"  
            },  
            "author": "Rita Liao",  
            "title": "Kai-Fu Lee's Sinovation bets on Linux tablet maker Jingling i...",  
            "description": "Kai-Fu Lee's Sinovation Ventures has its eyes on a nich...",  
            "url": "http://techcrunch.com/2021/06/15/jingos-10-million-linux-tablets-...",  
            "urlToImage": "https://techcrunch.com/wp-content/uploads/2021/06/Screen-S...",  
            "publishedAt": "2021-06-15T11:51:45Z",  
            "content": "Kai-Fu Lee's Sinovation Ventures has its eyes on a niche ma...",  
        },  
        {  
            "source": {  
                "id": "the-verge",  
                "name": "The Verge"  
            },  
            "author": "Sean Hollister",  
            "title": "AMD confirms it's powering the gaming rig inside Tesla's Mo...",  
            "description": "During its Computex 2021 keynote, AMD revealed that the n...",  
            "url": "https://www.theverge.com/2021/6/1/22462660/amd-tesla-model-x-s-pl...",  
            "urlToImage": "https://cdn.vox-cdn.com/thumbor/DLTaI4hph5QMrqGThuada3rxkI...",  
            "publishedAt": "2021-06-01T06:53:20Z",  
            "content": "Remember when Elon Musk claimed you'd be able to play The W...",  
        },  
        ... with 236 more lines  
    ]  
}
```

You can get your own API key at [News API's website](#).

Streaming APIs

Some web APIs return data in a streaming manner. This means that once you connect to the API, the data will continue to pour in until the connection is closed. A well-known example is the Twitter “fire hose,” which constantly streams all the tweets being sent around the world. Luckily, most command-line tools also operate in a streaming manner.

Let’s take a 10-second sample of one of Wikimedia’s streaming APIs, for example:

```
$ curl -s "https://stream.wikimedia.org/v2/stream/recentchange" |  
> sample -s 10 > wikimedia-stream-sample
```

This particular API returns all changes that have been made to Wikipedia and other properties of Wikimedia. The command-line tool `sample` is used to close the connection after 10 seconds. The connection can also be closed manually by pressing Ctrl-C to send an interrupt. The output is saved to the file `wikimedia-stream-sample`. Let’s take a peek using `trim`:

```
$ < wikimedia-stream-sample trim  
:ok  
  
event: message  
id:  
[{"topic":"eqiad.mediawiki.recentchange","partition":0,"timestamp":16101133...  
data: {"$schema":"/mediawiki/recentchange/1.0.0","meta":  
{"uri":"https://en.wiki...  
  
event: message  
id:  
[{"topic":"eqiad.mediawiki.recentchange","partition":0,"timestamp":16101133...  
data: {"$schema":"/mediawiki/recentchange/1.0.0","meta":  
{"uri":"https://www.wik...  
  
... with 1078 more lines
```

With a little bit of `sed` and `jq`, I can scrub this data to get a glimpse of the changes happening on the English version of Wikipedia:

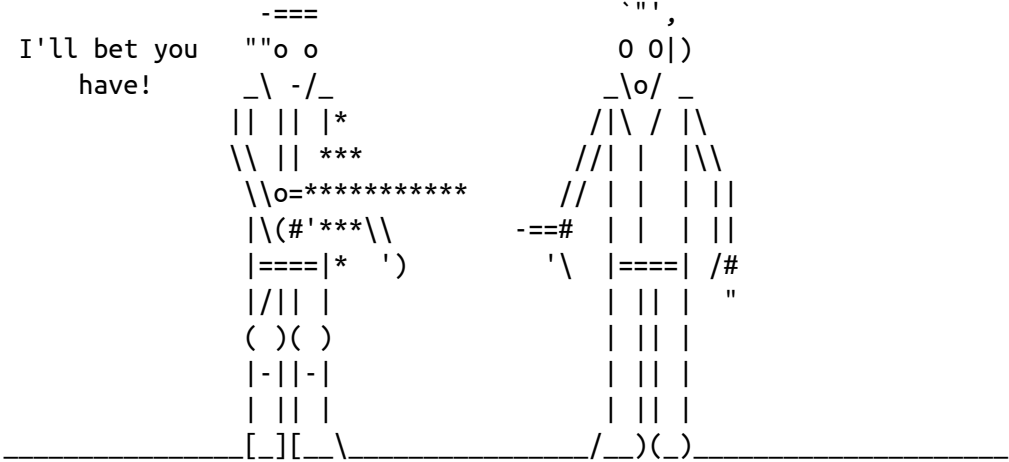
```
$ < wikipedia-stream-sample sed -n 's/^data: //p' | ❶
> jq 'select(.type == "edit" and .server_name == "en.wikipedia.org") | .title'
❷
"Odion Ighalo"
"Hold Up (song)"
"Talk:Royal Bermuda Yacht Club"
"Jenna Ushkowitz"
"List of films released by Yash Raj Films"
"SP.A"
"Odette (musician)"
"Talk:Pierre Avoi"
"User:Curlymanjaro/sandbox3"
"List of countries by electrification rate"
"Grieg (crater)"
"Gorman, Edmonton"
"Khabza"
"QAnon"
"Khaw Boon Wan"
"Draft:Oggy and the Cockroaches (1975 TV series)"
"Renzo Reggiardo"
"Greer, Arizona"
"National Curriculum for England"
"Mod DB"
"Jordanian Pro League"
"List of foreign Serie A players"
```

- ❶ This `sed` expression only prints lines that start with `data:` and prints the part after the semicolon, which happens to be JSON.
- ❷ This `jq` expression prints the `title` key of JSON objects that have a certain `type` and `server_name`.

Speaking of streaming, did you know that you could stream *Star Wars: Episode IV—A New Hope* for free using `telnet`?¹²

```
$ telnet towel.blinkenlights.nl
```

And after some time, we see that Han Solo did shoot first!



Sure, it's probably not a good source of data, but there's nothing wrong with enjoying an old classic while training your machine learning models.¹³

Summary

Congratulations, you have finished the first step of the OSEMN model. You've learned a variety of ways to obtain data, ranging from downloading to querying a relational database. In the next chapter, which is an intermezzo chapter, I'll teach you how to create your own command-line tools. But feel free to skip that discussion and go on to [Chapter 5](#) (the second step of the OSEMN model) if you cannot wait to learn about scrubbing data.

For Further Exploration

- Looking for a dataset to practice on? The GitHub repository *Awesome Public Datasets* lists hundreds of high-quality datasets that are publicly available.
 - Or perhaps you'd rather practice with an API: the GitHub repository *Public APIs* lists many free APIs. **City Bikes** and **The One API** are among my favorites.

- Writing SQL queries to obtain data from a relational database is an important skill. The first 15 lessons of the book *Sams Teach Yourself SQL in 10 Minutes a Day* by Ben Forta (Sams) teach the SELECT statement and its filtering, grouping, and sorting capabilities.
-

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- 12 Mats Erik Andersson, Andreas Henriksson, and Christoph Biedl, *telnet – User Interface to the TELNET Protocol*, version 0.17, 1999, <http://www.hcs.harvard.edu/~dholland/computers/netkit.html>.
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Chapter 4. Creating Command-Line Tools

Throughout the book, I'll introduce you to many commands and combinations of commands that basically fit on one line. These are known as *one-liners* or *pipelines*. Being able to perform complex tasks with just a one-liner is what makes the command line powerful. It's a very different experience from writing and using traditional programs.

Some tasks you perform only once, and some you perform more often. Some tasks are very specific, while others can be generalized. If you need to repeat a certain one-liner on a regular basis, it's worthwhile to turn this into a command-line tool of its own. So both one-liners and command-line tools have their uses. Recognizing the opportunity to turn a one-liner or existing code into a command-line tool requires practice and skill. The advantages of a command-line tool are that you don't have to remember the entire one-liner and that it improves readability if you include it into some other pipeline. In that sense, you can think of a command-line tool as similar to a function in a programming language.

The benefit of working with a programming language, however, is that the code is in one or more files. This means that you can easily edit and reuse that code. If the code has parameters, it can even be generalized and reapplied to problems that follow a similar pattern.

Command-line tools have the best of both worlds: they can be used from the command line, they accept parameters, and they have to be created only once. In this chapter, you're going to get familiar with creating command-line tools in two ways. First, I explain how to turn those one-liners into reusable command-line tools. By adding parameters to your commands, you can add the same flexibility that a programming language offers. Subsequently, I demonstrate how to create reusable command-line tools from code that's written in a programming language. By following the Unix

philosophy, your code can be combined with other command-line tools, which may be written in an entirely different language. In this chapter, I will focus on three programming languages: Bash, Python, and R.

I believe that creating reusable command-line tools makes you a more efficient and productive data scientist in the long run. You will gradually build up your own data science toolbox, from which you can draw existing tools and apply them to problems you have encountered previously.

TIP

To turn a one-liner into a shell script, I'm going to use a tiny bit of shell scripting. This book demonstrates only a small subset of concepts from shell scripting, including variables, conditionals, and loops. A complete course in shell scripting could fill a book all on its own and is therefore beyond the scope of this one. If you want to dive more deeply into shell scripting, I recommend the book *Classic Shell Scripting* by Arnold Robbins and Nelson H. F. Beebe (O'Reilly).

Overview

In this chapter, you'll learn how to:

- Convert one-liners into parameterized shell scripts
- Turn existing Python and R code into reusable command-line tools

This chapter starts with the following files:

```
$ cd /data/ch04

$ l
total 32K
-rwxr--r-- 1 dst dst 400 Jun 29 14:27 fizzbuzz.py*
-rwxr--r-- 1 dst dst 391 Jun 29 14:27 fizzbuzz.R*
-rwxr--r-- 1 dst dst 182 Jun 29 14:27 stream.py*
-rwxr--r-- 1 dst dst 147 Jun 29 14:27 stream.R*
-rwxr--r-- 1 dst dst 105 Jun 29 14:27 top-words-4.sh*
-rwxr--r-- 1 dst dst 128 Jun 29 14:27 top-words-5.sh*
-rwxr--r-- 1 dst dst 647 Jun 29 14:27 top-words.py*
-rwxr--r-- 1 dst dst 584 Jun 29 14:27 top-words.R*
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Converting One-Liners into Shell Scripts

In this section I'm going to explain how to turn a one-liner into a reusable command-line tool. Let's say that you would like to get the top 10 most frequently used words in a piece of text. Take the book *Alice's Adventures in Wonderland* by Lewis Carroll, which like many other great books, is freely available on Project Gutenberg:

```
$ curl -sL "https://www.gutenberg.org/files/11/11-0.txt" | trim  
The Project Gutenberg eBook of Alice's Adventures in Wonderland, by Lewis...
```

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... with 3751 more lines

The following sequence of tools, or *pipeline*, should do the job:

```
$ curl -sL "https://www.gutenberg.org/files/11/11-0.txt" | ❶  
> tr '[:upper:]' '[:lower:]' | ❷  
> grep -oE "[a-z']{2,}" | ❸  
> sort | ❹  
> uniq -c | ❺  
> sort -nr | ❻  
> head -n 10 ❻  
1839 the  
942 and  
811 to  
638 of  
610 it  
553 she  
486 you  
462 said
```

```
435 in  
403 alice
```

- ❶ Download an ebook using `curl`.
- ❷ Convert the entire text to lowercase using `tr`.¹
- ❸ Extract all the words using `grep`² and put each word on a separate line.
- ❹ Sort these words in alphabetical order using `sort`.
- ❺ Remove all the duplicates and count how often each word appears in the list using `uniq`.³
- ❻ Sort this list of unique words by their count in descending order using `sort`.
- ❼ Keep only the top 10 lines (i.e., words) using `head`.

Those words indeed appear the most often in the text. Because those words (apart from *alice*) appear very frequently in many English texts, they carry very little meaning. In fact, they are known as *stopwords*. If we get rid of those, we keep the most frequent words that are related to this text.

Here's a list of stopwords I've found:

```
$ curl -sL "https://raw.githubusercontent.com/stopwords-iso/stopwords-  
en/master/  
stopwords-en.txt" |  
> sort | tee stopwords | trim 20  
10  
39  
a  
able  
ableabout  
about
```

```
above
abroad
abst
accordance
according
accordingly
across
act
actually
ad
added
adj
adopted
ae
... with 1278 more lines
```

With `grep` we can filter out the stopwords right before we start counting:

```
$ curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
> tr '[:upper:]' '[:lower:]' |
> grep -oE "[a-z']{2,}" |
> sort |
> grep -Fwf stopwords | ❶
> uniq -c |
> sort -nr |
> head -n 10
 403 alice
   98 gutenberg
   88 project
   76 queen
   71 time
   63 king
   60 turtle
   57 mock
   56 hatter
   55 gryphon
```

- ❶ Obtain the patterns from a file (*stopwords* in our case), one pattern per line, with `-f`. Interpret those patterns as fixed strings with `-F`. Select only those lines containing matches that form whole words with `-w`. Select nonmatching lines with `-v`.

TIP

Each command-line tool used in this one-liner offers a man page. So in case you would like to know more about, say, `grep`, you can run `man grep` from the command line. The command-line tools `tr`, `grep`, `uniq`, and `sort` will be discussed in more detail in the next chapter.

There is nothing wrong with running this one-liner just once. However, imagine if you wanted to have the top 10 words of every ebook on Project Gutenberg. Or imagine that you wanted the top 10 words appearing on a news website on an hourly basis. In those cases, it would be best to have this one-liner as a separate building block that can be part of something bigger. To add some flexibility to this one-liner in terms of parameters, let's turn it into a shell script.

This allows us to take the one-liner as the starting point and gradually improve on it. To turn this one-liner into a reusable command-line tool, I'll walk you through the following six steps:

1. Copy and paste the one-liner into a file.
2. Add execute permissions.
3. Define a so-called shebang.
4. Remove the fixed input part.
5. Add a parameter.
6. Optionally extend your PATH.

Step 1: Create a File

The first step is to create a new file. You can open your favorite text editor and copy and paste the one-liner. Let's name the file `top-words-1.sh` to indicate that this is the first step towards our new command-line tool. If you like to stay at the command line, you can use the builtin `fc`, which stands for *fix command*, and allows you to fix or *edit* the last-run command:

```
$ fc
```

Running `fc` invokes the default text editor, which is stored in the environment variable `EDITOR`. In the Docker container, this is set to `nano`,⁴ a straightforward text editor. As you can see, this file contains our one-liner:

```
GNU nano 5.4 /tmp/zsh9198lv
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
tr '[:upper:]' '[:lower:]' |
grep -oE "[a-z\\']{2,}" |
sort |
grep -Fwf stopwords |
uniq -c |
sort -nr |
head -n 10
```

```
[ Read 8 lines ]
^G Help      ^O Write Out ^W Where Is  ^K Cut      ^T Execute  ^C Location
^X Exit      ^R Read File ^\ Replace   ^U Paste    ^J Justify  ^_ Go To Line
```

Let's give this temporary file a proper name by pressing `Ctrl-O`, removing the temporary filename, and typing `top-words-1.sh`:

```
GNU nano 5.4 /tmp/zsh9198lv
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
tr '[:upper:]' '[:lower:]' |
grep -oE "[a-z\\']{2,}" |
sort |
grep -Fwf stopwords |
uniq -c |
sort -nr |
head -n 10
```

```
File Name to Write: top-words-1.sh
^G Help      M-D DOS Format      M-A Append      M-B Backup File
^C Cancel    M-M Mac Format     M-P Prepend    ^T Browse
```

Press Enter:

```
GNU nano 5.4                               /tmp/zsh9198lv
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
tr '[:upper:]' '[:lower:]' |
grep -oE "[a-z\\']{2,}" |
sort |
grep -Fwf stopwords |
uniq -c |
sort -nr |
head -n 10
```

Save file under DIFFERENT NAME?

Y Yes
 N No Cancel

Press Y to confirm that you want to save under a different filename:

```
GNU nano 5.4                               top-words-1.sh
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
tr '[:upper:]' '[:lower:]' |
grep -oE "[a-z\\']{2,}" |
sort |
grep -Fwf stopwords |
uniq -c |
sort -nr |
head -n 10
```

[Wrote 8 lines]
^G Help ^O Write Out ^W Where Is ^K Cut ^T Execute ^C Location
^X Exit ^R Read File ^\ Replace ^U Paste ^J Justify ^_ Go To Line

Press Ctrl-X to exit **nano** and go back from whence you came.

We are using the file extension **.sh** to make clear that we are creating a shell script. However, command-line tools don't need to have an extension. In fact, command-line tools rarely have extensions.

Confirm the contents of the file:

```
$ pwd
/data/ch04

$ l
total 44K
-rwxr--r-- 1 dst dst  400 Jun 29 14:27 fizzbuzz.py*
-rwxr--r-- 1 dst dst  391 Jun 29 14:27 fizzbuzz.R*
-rw-r--r-- 1 dst dst 7.5K Jun 29 14:27 stopwords
-rwxr--r-- 1 dst dst 182 Jun 29 14:27 stream.py*
-rwxr--r-- 1 dst dst 147 Jun 29 14:27 stream.R*
-rw-r--r-- 1 dst dst 173 Jun 29 14:27 top-words-1.sh
-rwxr--r-- 1 dst dst 105 Jun 29 14:27 top-words-4.sh*
-rwxr--r-- 1 dst dst 128 Jun 29 14:27 top-words-5.sh*
-rwxr--r-- 1 dst dst 647 Jun 29 14:27 top-words.py*
-rwxr--r-- 1 dst dst 584 Jun 29 14:27 top-words.R*

$ bat top-words-1.sh
-----
| File: top-words-1.sh
-----
|-----|
| 1 | curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
| 2 | tr '[:upper:]' '[:lower:]' |
| 3 | grep -oE "[a-z\\']{2,}" |
| 4 | sort |
| 5 | grep -Fwf stopwords |
| 6 | uniq -c |
| 7 | sort -nr |
| 8 | head -n 10
|-----|
|-----|
```

You can now use `bash5` to interpret and execute the commands in the file:

```
$ bash top-words-1.sh
403 alice
98 gutenberg
88 project
76 queen
71 time
63 king
60 turtle
57 mock
```

```
56 hatter  
55 gryphon
```

This saves you from typing the one-liner again next time.

However, because the file cannot be executed on its own, it's not yet a *real* command-line tool. Let's change that in the next step.

Step 2: Give Permission to Execute

The reason we cannot execute our file directly is that we don't have the correct access permissions. In particular, you, as a user, need to have permission to execute the file. In this section we change the access permissions of our file.

In order to compare differences between steps, copy the file to *top-words-2.sh* using `cp -v top-words-{1,2}.sh`.

TIP

If you ever want to verify what the brace expansion or any other form of file expansion leads to, replace the command with `echo` to just print the result—for example, `echo book_{draft,final}.md` or `echo agent-{001..007}`.

To change the access permissions of a file, we need to use a command-line tool called `chmod`,⁶ which stands for *change mode*. It changes the file mode bits of a specific file. The following command gives the user (you), permission to execute *top-words-2.sh*:

```
$ cp -v top-words-{1,2}.sh  
'top-words-1.sh' -> 'top-words-2.sh'  
  
$ chmod u+x top-words-2.sh
```

The argument `u+x` consists of three characters: (1) `u` indicates that we want to change the permissions for the user who owns the file, which is you,

because you created the file; (2) + indicates that we want to add a permission; and (3) x indicates the permissions to execute.

Now let's have a look at the access permissions of both files:

```
$ ls top-words-{1,2}.sh
-rw-r--r-- 1 dst dst 173 Jun 29 14:27 top-words-1.sh
-rwxr--r-- 1 dst dst 173 Jun 29 14:28 top-words-2.sh*
```

The first column shows the access permissions for each file. For *top-words-2.sh*, this is *-rwxr--r--*. The first character, – (hyphen), indicates the file type. A – means regular file and a d means directory. The next three characters, *rwx*, indicate the access permissions for the user who owns the file. The *x* and *w* mean *read* and *write*, respectively. (As you can see, *top-words-1.sh* has a – instead of an *x*, which means that we cannot *execute* that file.) The next three characters, *rw-*, indicate the access permissions for all members of the group that owns the file. Finally, the last three characters in the column, *r--*, indicate access permissions for all other users.

Now you can execute the file as follows:

```
$ ./top-words-2.sh
403 alice
98 gutenberg
88 project
76 queen
71 time
63 king
60 turtle
57 mock
56 hatter
55 gryphon
```

If you try to execute a file for which you don't have the correct access permissions, as with *top-words-1.sh*, you will see the following error message:

```
$ ./top-words-1.sh  
zsh: permission denied: ./top-words-1.sh
```

Step 3: Define a Shebang

Although we can already execute the file on its own, we should add a so-called shebang to the file. The *shebang* is a special line in the script that instructs the system as to which executable it should use to interpret the commands.

The name *shebang* comes from the first two characters: a hash (she) and an exclamation mark (bang): `#!`. It's not a good idea to leave it out like we have done in the previous step, because each shell has a different default executable. The Z shell, the one we're using throughout the book, uses the executable `/bin/sh` by default if no shebang is defined. In this case I'd like `bash` to interpret the commands, as that will give us some more functionality than `sh` would.

Again, you're free to use whatever editor you like, but I'm going to stick with `nano`, which is installed in the Docker image:

```
$ cp -v top-words-{2,3}.sh  
'top-words-2.sh' -> 'top-words-3.sh'
```

```
$ nano top-words-3.sh
```

```
GNU nano 5.4                               top-words-3.sh  
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |  
tr '[:upper:]' '[:lower:]' |  
grep -oE "[a-z\\']{2,}" |  
sort |  
grep -Fwf stopwords |  
uniq -c |  
sort -nr |  
head -n 10
```

[Read 8 lines]

<code>^G</code> Help	<code>^O</code> Write Out	<code>^W</code> Where Is	<code>^K</code> Cut	<code>^T</code> Execute	<code>^C</code> Location
<code>^X</code> Exit	<code>^R</code> Read File	<code>^\\</code> Replace	<code>^U</code> Paste	<code>^J</code> Justify	<code>^_</code> Go To Line

Go ahead and type `#!/usr/bin/env bash` and press Enter. When you're ready, press Ctrl-X to save and exit:

```
GNU nano 5.4                               top-words-3.sh *
#!/usr/bin/env bash
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
tr '[:upper:]' '[:lower:]' |
grep -oE "[a-z\\']{2,}" |
sort |
grep -Fwf stopwords |
uniq -c |
sort -nr |
head -n 10
```

Save modified buffer?

Y Yes
 N No ^C Cancel

Press Y to indicate that you want to save the file:

```
GNU nano 5.4                               top-words-3.sh *
#!/usr/bin/env bash
curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
tr '[:upper:]' '[:lower:]' |
grep -oE "[a-z\\']{2,}" |
sort |
grep -Fwf stopwords |
uniq -c |
sort -nr |
head -n 10
```

File Name to Write: top-words-3.sh

^G Help M-D DOS Format M-A Append M-B Backup File
 ^C Cancel M-M Mac Format M-P Prepend ^T Browse

Let's confirm what `top-words-3.sh` looks like:

```
$ bat top-words-3.sh
```

```
| File: top-words-3.sh
|
1 | #!/usr/bin/env bash
2 | curl -sL "https://www.gutenberg.org/files/11/11-0.txt" |
3 | tr '[:upper:]' '[:lower:]' |
4 | grep -oE "[a-z']{2,}" |
5 | sort |
6 | grep -Fwf stopwords |
7 | uniq -c |
8 | sort -nr |
9 | head -n 10
```

That's exactly what we need: our original pipeline with a shebang in front of it.

Sometimes you will come across scripts that have a shebang in the form of `!/usr/bin/bash`, or `!/usr/bin/python` in the case of Python (as we will see in the next section). While this generally works, if the `bash` or `python`⁷ executables are installed in a different location than `/usr/bin`, then the script does not work anymore. It is better to use the form that I present here, namely `!/usr/bin/env bash` and `!/usr/bin/env python`, because the `env`⁸ executable is aware of where `bash` and `python` are installed. In short, using `env` makes your scripts more portable.

Step 4: Remove the Fixed Input

We now have a valid command-line tool that we can execute from the command line. But we can do even better. We can make our command-line tool more reusable. The first command in our file is `curl`, which downloads the text from which we wish to obtain the top 10 most-used words. So the data and operations are combined into one.

What if we wanted to obtain the top 10 most-used words from another ebook, or from any other text for that matter? The input data is fixed within the tools itself. It would be better to separate the data from the command-line tool.

If we assume that the user of the command-line tool will provide the text, the tool will become generally applicable. So the solution is to remove the `curl` command from the script. Here is the updated script named *top-words-4.sh*:

```
$ cp -v top-words-{3,4}.sh  
'top-words-3.sh' -> 'top-words-4.sh'  
  
$ sed -i '2d' top-words-4.sh  
  
$ bat top-words-4.sh  
----  
|  
| File: top-words-4.sh  
|  
|  
| 1 | #!/usr/bin/env bash  
| 2 | tr '[:upper:]' '[:lower:]'  
| 3 | grep -oE "[a-z']{2,}" |  
| 4 | sort |  
| 5 | grep -Fwf stopwords |  
| 6 | uniq -c |  
| 7 | sort -nr |  
| 8 | head -n 10  
|
```

This works because if a script starts with a command that needs data from standard input, like `tr`, it will take the input that is given to the command-line tools. For example:

```
$ curl -sL 'https://www.gutenberg.org/files/11/11-0.txt' | ./top-words-4.sh
403 alice
98 gutenberg
88 project
76 queen
71 time
63 king
60 turtle
57 mock
56 hatter
55 gryphon
```

```
469 alice
189 queen
 98 gutenberg
 88 project
 72 time
 71 red
 70 white
 67 king
 63 head
 59 knight

$ man bash | ./top-words-4.sh
585 command
332 set
313 word
313 option
304 file
300 variable
298 bash
258 list
257 expansion
238 history
```

TIP

Although we have not done so in our script, the same principle holds for saving data. In general, it is better to let the user take care of that using output redirection than to let the script write to a specific file. Of course, if you intend to use a command-line tool only for your own projects, then there are no limits to how specific you can be.

Step 5: Add Arguments

There is one more step to making our command-line tool even more reusable: parameters. In our command-line tool, there are a number of fixed command-line arguments—for example, `-nr` for `sort` and `-n 10` for `head`. It is probably best to keep the former argument fixed. However, it would be very useful to allow for different values for the `head` command. This would allow the end user to set the number of most-often-used words to output. The following shows what our file `top-words-5.sh` looks like:

```
$ bat top-words-5.sh
```

```
File: top-words-5.sh
```

```
1  #!/usr/bin/env bash
2
3  NUM_WORDS="${1:-10}"
4
5  tr '[:upper:]' '[:lower:]' |
6  grep -oE "[a-z']{2,}" |
7  sort |
8  grep -Fwf stopwords |
9  uniq -c |
10 sort -nr |
11 head -n "${NUM_WORDS}"
```

- The variable `NUM_WORDS` is set to the value of `$1`, which is a special variable in Bash; it holds the value of the first command-line argument passed to our command-line tool. The table below lists the other special variables that Bash offers. If no value is specified, `NUM_WORDS` will take on the value `10`.
- Note that to *use* the value of the `NUM_WORDS` variable, you need to put a dollar sign in front of it. When you *set* it, you don't write a dollar sign.

We could have used `$1` directly as an argument for `head` and not bothered creating an extra variable such as `NUM_WORDS`. However, with larger scripts and a few more command-line arguments such as `$2` and `$3`, your code becomes more readable when you use named variables.

Now if we wanted to see the top 20 most-used words of our text, we would invoke our command-line tool as follows:

```
$ curl -sL "https://www.gutenberg.org/files/11/11-0.txt" > alice.txt

$ < alice.txt ./top-words-5.sh 20
403 alice
```

```
98 gutenberg
88 project
76 queen
71 time
63 king
60 turtle
57 mock
56 hatter
55 gryphon
53 rabbit
50 head
48 voice
45 looked
44 mouse
42 duchess
40 tone
40 dormouse
37 cat
34 march
```

If the user does not specify a number, then our script will show the top 10 most common words:

```
$ < alice.txt ./top-words-5.sh
403 alice
98 gutenberg
88 project
76 queen
71 time
63 king
60 turtle
57 mock
56 hatter
55 gryphon
```

Step 6: Extend Your PATH

After the previous five steps, we are finally finished building a reusable command-line tool. There is, however, one more step that can be very useful. In this optional step, we are going to ensure that you can execute your command-line tools from everywhere.

Currently, when you want to execute your command-line tool, you either have to navigate to the directory it is in or include the full pathname, as shown in step 2. This is fine if the command-line tool is built specifically for a certain project. However, if your command-line tool could be applied in multiple situations, then it is useful to be able to execute it from everywhere, just like the command-line tools that come with Ubuntu.

To accomplish this, Bash needs to know where to look for your command-line tools. It does this by traversing a list of directories that are stored in an environment variable called `PATH`. In a fresh Docker container, the `PATH` looks like this:

```
$ echo $PATH  
/usr/local/lib/R/site-  
library/rush/exec:/usr/bin/dsutils:/home/dst/.local/bin:/u  
sr/local/sbin:/usr/local/bin:/usr/sbin:/usr/bin:/sbin:/bin
```

The directories are delimited by colons. We can print this as a list of directories by *translating* the colons to newlines:

```
$ echo $PATH | tr ':' '\n'  
/usr/local/lib/R/site-library/rush/exec  
/usr/bin/dsutils  
/home/dst/.local/bin  
/usr/local/sbin  
/usr/local/bin  
/usr/sbin  
/usr/bin  
/sbin  
/bin
```

To change the `PATH` permanently, you'll need to edit the `.bashrc` or `.profile` file located in your home directory. If you put all your custom command-line tools into one directory—say, `~/tools`—then you only change the `PATH` once. Now you no longer need to add the `./` and can just use the filename. Moreover, you no longer need to remember where the command-line tool is located:

```

$ cp -v top-words{-5.sh,}
'top-words-5.sh' -> 'top-words'

$ export PATH="${PATH}:/data/ch04"

$ echo $PATH
/usr/local/lib/R/site-
library/rush/exec:/usr/bin/dsutils:/home/dst/.local/bin:/u
sr/local/sbin:/usr/local/bin:/usr/sbin:/usr/bin:/sbin:/bin:/data/ch04

$ curl "https://www.gutenberg.org/files/11/11-0.txt" |
> top-words 10
% Total    % Received % Xferd  Average Speed   Time     Time     Time
Current                                         Dload  Upload   Total Spent   Left  Speed
100  170k  100  170k    0      0  223k      0 --::-- --::-- --::-- 223k
 403 alice
  98 gutenberg
  88 project
  76 queen
  71 time
  63 king
  60 turtle
  57 mock
  56 hatter
  55 gryphon

```

Creating Command-Line Tools with Python and R

The command-line tool that we created in the previous section was written in Bash. (Sure, not every feature of the Bash programming language was employed, but the interpreter still was `bash`.) As you know by now, the command line is language agnostic, so we don't necessarily have to use Bash for creating command-line tools.

In this section I'm going demonstrate that command-line tools can be created in other programming languages as well. I'll focus on Python and R because these are the two most popular programming languages within the data science community. I cannot offer a complete introduction to either language, so I assume that you have some familiarity with Python and/or R.

Other programming languages such as Java, Go, and Julia follow a similar pattern when it comes to creating command-line tools.

There are three main reasons for creating command-line tools in a programming language other than Bash. First, you may already have some code that you'd like to be able to use from the command line. Second, the command-line tool would end up encompassing more than a hundred lines of Bash code. Third, the command-line tool needs to be more safe and robust (Bash lacks many features, such as type checking).

The six steps that I discussed in the previous section roughly apply to creating command-line tools in other programming languages as well. The first step, however, would not be copy and pasting from the command line; rather, it would be copy and pasting the relevant code into a new file.

Command-line tools written in Python and R need to specify `python` and `Rscript`,⁹ respectively, as the interpreter after the shebang.

When it comes to creating command-line tools using Python and R, there are two more aspects that deserve special attention. First, processing standard input, which comes naturally to shell scripts, has to be taken care of explicitly in Python and R. Second, as command-line tools written in Python and R tend to be more complex, we may also want to offer the user the ability to specify more elaborate command-line arguments.

Porting the Shell Script

As a starting point, let's see how we would port the shell script we just created to both Python and R. In other words, what Python and R code gives us the top most-often-used words from standard input? We will first show the two files `top-words.py` and `top-words.R` and then discuss the differences with the shell code. In Python, the code would look something like the following:

```
$ cd /data/ch04  
$ bat top-words.py
```

| File: **top-words.py**

```
1  #!/usr/bin/env python
2  import re
3  import sys
4
5  from collections import Counter
6  from urllib.request import urlopen
7
8  def top_words(text, n):
9      with urlopen("https://raw.githubusercontent.com/stopwords-
10 iso/stopw
11 ords-en/master/stopwords-en.txt") as f:
12         stopwords = f.read().decode("utf-8").split("\n")
13
14         words = re.findall("[a-z']{2,}", text.lower())
15         words = (w for w in words if w not in stopwords)
16
17         for word, count in Counter(words).most_common(n):
18             print(f"{count:>7} {word}")
19
20     if __name__ == "__main__":
21         text = sys.stdin.read()
22
23         try:
24             n = int(sys.argv[1])
25         except:
26             n = 10
27
28     top_words(text, n)
```

Note that this Python example doesn't use any third-party packages. If you want to do advanced text processing, then I recommend you check out the NLTK package.¹⁰ If you're going to work with a lot of numerical data, then I recommend you use the Pandas package.¹¹

In R the code would look something like this:

\$ bat **top-words.R**

| File: **top-words.R**

```

1 | #!/usr/bin/env Rscript
2 | n <- as.integer(commandArgs(trailingOnly = TRUE))
3 | if (length(n) == 0) n <- 10
4 |
5 | f_stopwords <- url("https://raw.githubusercontent.com/stopwords-
iso/sto
      pwords-en/master/stopwords-en.txt")
6 | stopwords <- readLines(f_stopwords, warn = FALSE)
7 | close(f_stopwords)
8 |
9 | f_text <- file("stdin")
10 | lines <- tolower(readLines(f_text))
11 |
12 | words <- unlist(regmatches(lines, gregexpr("[a-z'][{2,}", lines)))
13 | words <- words[is.na(match(words, stopwords))]
14 |
15 | counts <- sort(table(words), decreasing = TRUE)
16 | cat(sprintf("%7d %s\n", counts[1:n], names(counts[1:n])), sep = ""))
17 | close(f_text)

```

Let's check that all three implementations (i.e., Bash, Python, and R) return the same top five words with the same counts:

```

$ time < alice.txt top-words 5
403 alice
98 gutenberg
88 project
76 queen
71 time
top-words 5 < alice.txt 0.08s user 0.01s system 107% cpu 0.084 total

$ time < alice.txt top-words.py 5
403 alice
98 gutenberg
88 project
76 queen
71 time
top-words.py 5 < alice.txt 0.38s user 0.02s system 82% cpu 0.478 total

$ time < alice.txt top-words.R 5
403 alice

```

```
98 gutenberg
88 project
76 queen
71 time
top-words.R 5 < alice.txt  0.29s user 0.07s system 56% cpu 0.652 total
```

Wonderful! Sure, the output itself is not very exciting. What’s exciting is that we can accomplish the same task with multiple languages. Let’s look at the differences between the approaches.

First, what’s immediately obvious are the differences in the amount of code. For this specific task, both Python and R require much more code than Bash. This illustrates that, for some tasks, it is better to use the command line. For other tasks, you may be better off using a programming language. As you gain more experience on the command line, you will start to recognize when to use which approach. When everything is a command-line tool, you can even split up the task into subtasks and combine a Bash command-line tool with, say, a Python command-line tool—whichever approach works best for the task at hand.

Processing Streaming Data from Standard Input

In the previous two code snippets, both Python and R read the complete standard input at once. On the command line, most tools pipe data to the next command-line tool in a streaming fashion. A few command-line tools such as `sort` require the complete data before they write any data to standard output. This means the pipeline is blocked by these tools. This doesn’t have to be a problem when the input data is finite, like a file. However, when the input data is a nonstop stream, such blocking command-line tools are useless.

Luckily, Python and R support processing streaming data. You can apply a function on a line-per-line basis, for example. Here are two minimal examples that demonstrate how this works in Python and R, respectively.

Both the Python and R tools solve the by-now-infamous Fizz Buzz problem, which is defined as follows: print every number from 1 to 100, but if the number is divisible by 3, print “fizz” instead; if the number is

divisible by 5, print “buzz”; and if the number is divisible by 15, print “fizzbuzz.” Here’s the Python code:¹²

```
$ bat fizzbuzz.py
-----
| File: fizzbuzz.py
-----
1 #!/usr/bin/env python
2 import sys
3
4 CYCLE_OF_15 = ["fizzbuzz", None, None, "fizz", None,
5                 "buzz", "fizz", None, None, "fizz",
6                 "buzz", None, "fizz", None, None]
7
8 def fizz_buzz(n: int) -> str:
9     return CYCLE_OF_15[n % 15] or str(n)
10
11 if __name__ == "__main__":
12     try:
13         while (n:= sys.stdin.readline()):
14             print(fizz_buzz(int(n)))
15     except:
16         pass
```

And here’s the R code:

```
$ bat fizzbuzz.R
-----
| File: fizzbuzz.R
-----
1 #!/usr/bin/env Rscript
2 cycle_of_15 <- c("fizzbuzz", NA, NA, "fizz", NA,
3                   "buzz", "fizz", NA, NA, "fizz",
4                   "buzz", NA, "fizz", NA, NA)
5
6 fizz_buzz <- function(n) {
7     word <- cycle_of_15[as.integer(n) %% 15 + 1]
8     ifelse(is.na(word), n, word)
9 }
```

```
10  |   f <- file("stdin")
11  |   open(f)
12  |   while(length(n <- readLines(f, n = 1)) > 0) {
13  |     write(fizz_buzz(n), stdout())
14  |   }
15  |   close(f)
```

—

Let's test both tools (to save space, I've piped the output to `column`):

```
$ seq 30 | fizzbuzz.py | column -x
1          2          fizz        4          buzz
fizz      7          8          fizz        buzz
11         fizz      13         14         fizzbuzz
16         17         fizz      19         buzz
fizz      22         23         fizz        buzz
26         fizz      28         29         fizzbuzz

$ seq 30 | fizzbuzz.R | column -x
1          2          fizz        4          buzz
fizz      7          8          fizz        buzz
11         fizz      13         14         fizzbuzz
16         17         fizz      19         buzz
fizz      22         23         fizz        buzz
26         fizz      28         29         fizzbuzz
```

This output looks correct to me! It's difficult to demonstrate that these two tools actually work in a streaming manner. You can verify this yourself by piping the input data to `sample -d 100` before it's piped to the Python or R tool. That way, you'll add a small delay in between each line so that it's easier to confirm that the tools don't wait for all the input data but instead operate on a line-by-line basis.

Summary

In this intermezzo chapter, I have shown you how to build your own command-line tool. Only six steps are needed to turn your code into a reusable building block, which you'll find makes you much more

productive. I advise you to keep an eye out for opportunities to create your own tools. The next chapter covers the second step of the OSEMN model for data science, namely scrubbing data.

For Further Exploration

- Adding help documentation to your tool becomes important when the tool has many options to remember, and even more so when you want to share your tool with others. `docopt` is a language-agnostic framework for providing help and defining the options that your tool accepts. Implementations are available in just about any programming language, including Bash, Python, and R.
- If you want to learn more about programming in Bash, I recommend *Classic Shell Programming* by Arnold Robbins and Nelson H. F. Beebe (O'Reilly) and *Bash Cookbook, 2nd Edition* by Carl Albing and JP Vossen (O'Reilly).
- Writing a robust and safe Bash script is quite tricky. `ShellCheck` is an online tool that will check your Bash code for mistakes and vulnerabilities. A command-line tool is also available.
- The book *Ten Essays on Fizz Buzz* by Joel Grus (Brightwalton) is an insightful and fun collection of 10 different ways to solve Fizz Buzz with Python.

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<https://www.gnu.org/software/grep>.

³ Richard M. Stallman and David MacKenzie, *uniq – Report or Omit Repeated Lines*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

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<https://nano-editor.org>.

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- 6 David MacKenzie and Jim Meyering, *chmod – Change File Mode Bits*, version 8.30, 2018, <https://www.gnu.org/software/coreutils>.
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- 8 Richard Mlynarik, David MacKenzie, and Assaf Gordon, *env – Run a Program in a Modified Environment*, version 8.32, 2020, <https://www.gnu.org/software/coreutils>.
- 9 The R Foundation for Statistical Computing, *R – a Language and Environment for Statistical Computing*, version 4.0.4, 2021, <https://www.r-project.org>.
- 10 Jacob Perkins, *Python Text Processing with NLTK 2.0 Cookbook* (Birmingham, UK: Packt, 2010).
- 11 Wes McKinney, *Python for Data Analysis* (O'Reilly, 2017).
- 12 This code is adapted from a [Python script by Joel Grus](#).

Chapter 5. Scrubbing Data

Two chapters ago, in the first step of the OSEMN model for data science, we looked at *obtaining* data from a variety of sources. This chapter is all about the second step: *scrubbing* data. You see, it's quite rare that you can move directly from obtaining data to *exploring* or even *modeling* the data. There's a plethora of reasons why your data first needs some cleaning, or scrubbing.

For starters, the data might not be in the desired format. For example, you may have obtained some JSON data from an API, but you need it to be in CSV format to create a visualization. Other common formats include plain text, HTML, and XML. Most command-line tools work with only one or two formats, so it's important that you're able to convert data from one format to another.

Once the data is in the desired format, there could still be issues like missing values, inconsistencies, weird characters, or unnecessary parts. You can fix these by applying filters, replacing values, and combining multiple files. The command line is especially well suited for these kind of transformations, because there are many specialized tools available, most of which can handle large amounts of data. In this chapter I'll discuss classic tools such as `grep` and `awk`,¹ and newer tools such as `jq` and `pup`.

Sometimes you can use the same command-line tool to perform several operations or multiple tools to perform the same operation. This chapter is structured more like a cookbook in that it focuses on the problems or recipes rather than diving deeply into the command-line tools themselves.

Overview

In this chapter, you'll learn how to:

- Convert data from one format to another

- Apply SQL queries directly to CSV
- Filter lines
- Extract and replace values
- Split, merge, and extract columns
- Combine multiple files

This chapter starts with the following files:

```
$ cd /data/ch05

$ l
total 200K
-rw-r--r-- 1 dst dst 164K Jun 29 14:28 alice.txt
-rw-r--r-- 1 dst dst 4.5K Jun 29 14:28 iris.csv
-rw-r--r-- 1 dst dst 179 Jun 29 14:28 irismeta.csv
-rw-r--r-- 1 dst dst 160 Jun 29 14:28 names-comma.csv
-rw-r--r-- 1 dst dst 129 Jun 29 14:28 names.csv
-rw-r--r-- 1 dst dst 7.8K Jun 29 14:28 tips.csv
-rw-r--r-- 1 dst dst 5.1K Jun 29 14:28 users.json
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Before I dive into the actual transformations, I'd like to demonstrate their ubiquity when working at the command line.

Transformations, Transformations Everywhere

In [Chapter 1](#) I mentioned that, in practice, the steps of the OSEMN model will rarely be followed linearly. In this vein, although scrubbing is the second step of the OSEMN model, I want you to know that it's not *just* the obtained data that needs scrubbing. The transformations that you'll learn in this chapter can be useful at any part of your pipeline and at any step of the OSEMN model. Generally, if one command-line tool generates output that

can be used immediately by the next tool, you can chain the two tools together by using the pipe operator (`|`). Otherwise, a transformation first needs to be applied to the data by inserting an intermediate tool into the pipeline.

Let me walk you through an example to make this more concrete. Imagine that you have obtained the first one hundred items of a Fizz Buzz sequence (“[Processing Streaming Data from Standard Input](#)”) and that you’d like to visualize how often the words *fizz*, *buzz*, and *fizzbuzz* appear using a bar chart. Don’t worry if this example uses tools that you might not be familiar with yet; they’ll all be covered in more detail later.

First you obtain the data by generating the sequence and write it to `fb.seq`:

```
$ seq 100 |  
> /data/ch04/fizzbuzz.py | ❶  
> tee fb.seq | trim  
1  
2  
fizz  
4  
buzz  
fizz  
7  
8  
fizz  
buzz  
... with 90 more lines
```

❶ The custom tool `fizzbuzz.py` comes from [Chapter 4](#).

Then you use `grep` to keep the lines that match the pattern *fizz* or *buzz* and count how often each word appears using `sort` and `uniq`:

```
$ grep -E "fizz|buzz" fb.seq | ❶  
> sort | uniq -c | sort -nr > fb.cnt ❷  
  
$ bat -A fb.cnt
```

```
| File: fb.cnt
-----
| 1 | .....27·fizz ↵
| 2 | .....14·buzz ↵
| 3 | .....6·fizzbuzz ↵
```

- ① This regular expression also matches *fizzbuzz*.
- ② Using `sort` and `uniq` like this is a common way to count lines and sort them in descending order. It's the `-c` option that adds the counts.

Note that `sort` is used twice: first because `uniq` assumes its input data to be sorted, and second to sort the counts numerically. In a way, this is an intermediate transformation, albeit a subtle one.

The next step would be to visualize the counts using `rush`.² However, since `rush` expects the input data to be in CSV format, this requires an initial, less subtle transformation. `awk` can add a header, flip the two fields, and insert commas in a single incantation:

```
$ < fb.cnt awk 'BEGIN { print "value,count" } { print $2","$1 }' > fb.csv
```

```
$ bat fb.csv
```

```
| File: fb.csv
```

```
1 | value,count
2 | fizz,27
3 | buzz,14
4 | fizzbuzz,6
```

```
$ csvlook fb.csv
| value      | count   |
```

fizz	27
buzz	14
fizzbuzz	6

Now you're ready to use `rush` to create a bar chart; see [Figure 5-1](#) for the result (I'll cover this syntax of `rush` in detail in [Chapter 7](#)):

```
$ rush plot -x value -y count --geom col --height 2 fb.csv > fb.png  
$ display fb.png
```

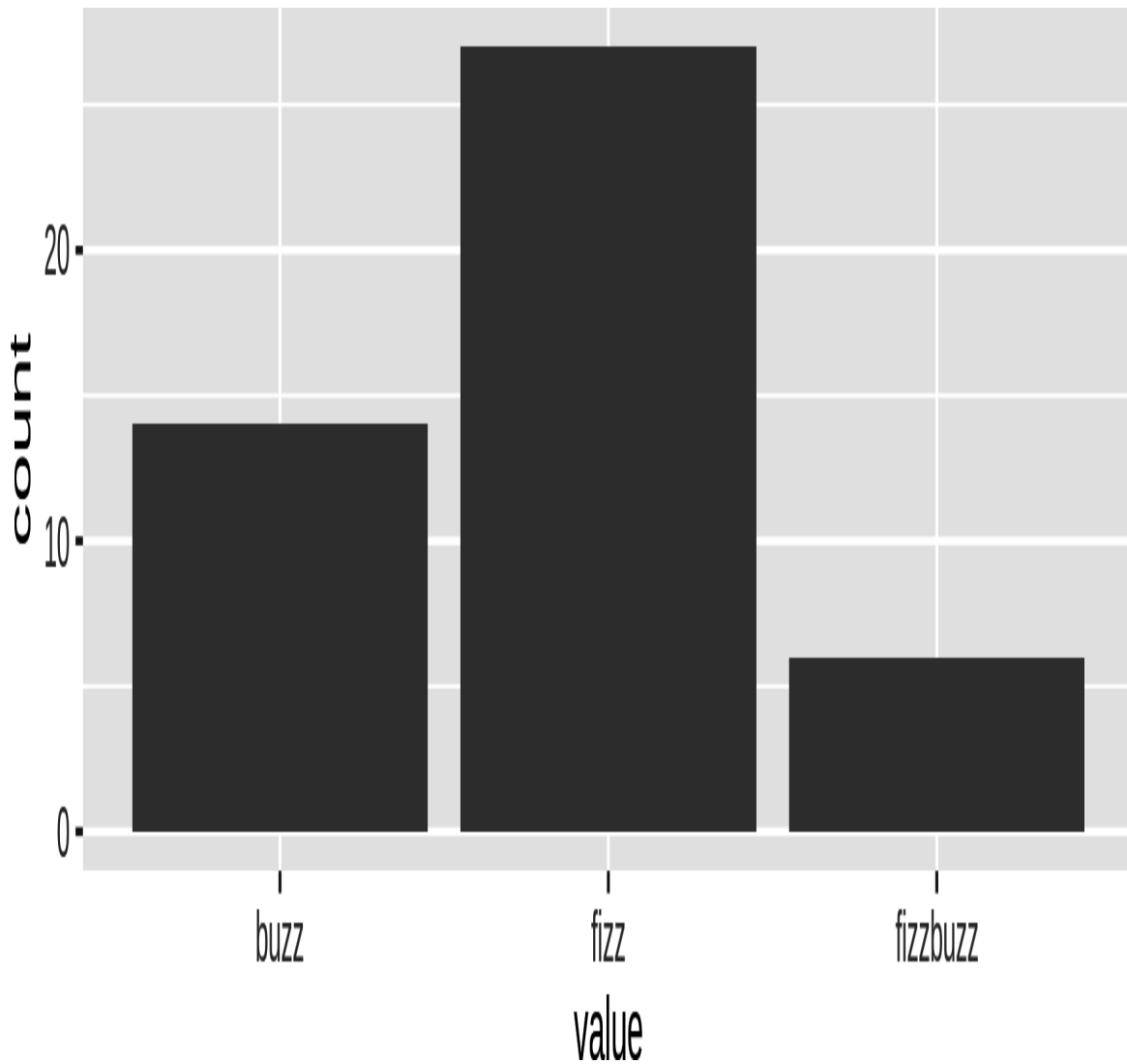


Figure 5-1. Counting fizz, buzz, and fizzbuzz

Although this example is a bit contrived, it reveals a pattern that is common when working at the command line. The key tools, such as the ones that obtain data, create a visualization, or train a model, often require intermediate transformations in order to be chained into a pipeline. In that sense, writing a pipeline is like solving a puzzle, where the key pieces often require helper pieces to fit.

Now that you've seen the importance of scrubbing data, you're ready to learn about some actual transformations.

Plain Text

Formally speaking, *plain text* refers to a sequence of human-readable characters and, optionally, some specific types of control characters such as tabs and newlines.³ Examples are logs, ebooks, emails, and source code. Plain text has many advantages over binary data, including the following:⁴

- It can be opened, edited, and saved using any text editor.
- It's self-describing and independent of the application that created it.
- It will outlive other forms of data, because no additional knowledge or applications are required to process it.

Most importantly, the Unix philosophy considers plain text to be the universal interface between command-line tools.⁵ This means that most tools accept plain text as input and produce plain text as output.

That's reason enough for me to start with plain text. The other formats that I discuss in this chapter—CSV, JSON, XML, and HTML—are indeed also plain text. For now, I assume that the plain text has no clear tabular structure (like CSV does) or nested structure (like JSON, XML, and HTML do). Later in this chapter, I'll introduce some tools that are specifically designed for working with these formats.

Filtering Lines

The first scrubbing operation is filtering lines. This means that each line from the input data will be evaluated on whether it will be kept or discarded.

Based on location

The most straightforward way to filter lines is based on their location. This may be useful when you want to inspect, say, the top 10 lines of a file, or when you extract a specific row from the output of another command-line tool. To illustrate how to filter based on location, let's create a file that contains 10 lines:

```
$ seq -f "Line %g" 10 | tee lines
Line 1
Line 2
Line 3
Line 4
Line 5
Line 6
Line 7
Line 8
Line 9
Line 10
```

You can print the first three lines using either `head`, `sed`,⁶ or `awk`:

```
$ < lines head -n 3
Line 1
Line 2
Line 3

$ < lines sed -n '1,3p'
Line 1
Line 2
Line 3

$ < lines awk 'NR <= 3' ①
Line 1
Line 2
Line 3
```

- ① In **awk**, *NR* refers to the total number of input records seen so far.

Similarly, you can print the last three lines using **tail**:⁷

```
$ < lines tail -n 3
Line 8
Line 9
Line 10
```

You can also use **sed** and **awk** for this, but **tail** is much faster. Removing the first three lines goes as follows:

```
$ < lines tail -n +4
Line 4
Line 5
Line 6
Line 7
Line 8
Line 9
Line 10

$ < lines sed '1,3d'
Line 4
Line 5
Line 6
Line 7
Line 8
Line 9
Line 10

$ < lines sed -n '1,3!p'
Line 4
Line 5
Line 6
Line 7
Line 8
Line 9
Line 10
```

Notice that with **tail** you have to specify the number of lines you want to remove plus one. Think of it as the line from which you want to start

printing. Removing the last three lines can be done with `head`:

```
$ < lines head -n -3
Line 1
Line 2
Line 3
Line 4
Line 5
Line 6
Line 7
```

You can print specific lines using either `sed`, `awk`, or a combination of `head` and `tail`. Here I print lines 4, 5, and 6:

```
$ < lines sed -n '4,6p'
Line 4
Line 5
Line 6

$ < lines awk '(NR>=4) && (NR<=6)'
Line 4
Line 5
Line 6

$ < lines head -n 6 | tail -n 3
Line 4
Line 5
Line 6
```

You can print odd lines with `sed` by specifying a start and a step, or with `awk` by using the modulo operator:

```
$ < lines sed -n '1~2p'
Line 1
Line 3
Line 5
Line 7
Line 9

$ < lines awk 'NR%2'
Line 1
Line 3
Line 5
```

```
Line 7  
Line 9
```

Printing even lines works in a similar manner:

```
$ < lines sed -n '0~2p'  
Line 2  
Line 4  
Line 6  
Line 8  
Line 10
```

```
$ < lines awk '(NR+1)%2'  
Line 2  
Line 4  
Line 6  
Line 8  
Line 10
```

NOTE

Many of these examples start with the less-than sign (<) followed by the filename. I do this because it allows me to read the pipeline from left to right. Please know that this is my own preference. You can also use `cat` to pipe the contents of a file. Additionally, many command-line tools accept the filename as an argument.

Based on a pattern

Sometimes you want to keep or discard lines based on their contents. With `grep`, the canonical command-line tool for filtering lines, you can print every line that matches a certain pattern or regular expression. For example, to extract all the chapter headings from *Alice's Adventures in Wonderland*:

```
$ < alice.txt grep -i chapter ❶  
CHAPTER I. Down the Rabbit-Hole  
CHAPTER II. The Pool of Tears  
CHAPTER III. A Caucus-Race and a Long Tale  
CHAPTER IV. The Rabbit Sends in a Little Bill  
CHAPTER V. Advice from a Caterpillar  
CHAPTER VI. Pig and Pepper  
CHAPTER VII. A Mad Tea-Party  
CHAPTER VIII. The Queen's Croquet-Ground
```

```
CHAPTER IX. The Mock Turtle's Story  
CHAPTER X. The Lobster Quadrille  
CHAPTER XI. Who Stole the Tarts?  
CHAPTER XII. Alice's Evidence
```

- ❶ The `-i` option specifies that the matching should be case insensitive.

You can also specify a regular expression. For example, if you wanted to print only the headings that start with *The*:

```
$ < alice.txt grep -E '^CHAPTER (.*)\\. The'  
CHAPTER II. The Pool of Tears  
CHAPTER IV. The Rabbit Sends in a Little Bill  
CHAPTER VIII. The Queen's Croquet-Ground  
CHAPTER IX. The Mock Turtle's Story  
CHAPTER X. The Lobster Quadrille
```

Note that you have to specify the `-E` option to enable regular expressions. Otherwise, `grep` interprets the pattern as a literal string, which most likely results in no matches at all:

```
$ < alice.txt grep '^CHAPTER (.*)\\. The'
```

With the `-v` option, you invert the matches, so that `grep` prints the lines that *don't* match the pattern. The regular expression below matches lines that contain white space only. So with the inverse, and using `wc -l`, you can count the number of non-empty lines:

```
$ < alice.txt grep -Ev '^\\s$' | wc -l  
2790
```

Based on randomness

When you're in the process of formulating your data pipeline and you have a lot of data, debugging your pipeline can be cumbersome. In that case, generating a smaller sample from the data might be useful. This is where `sample`⁸ comes in handy. The main purpose of `sample` is to get a subset of

the data by outputting only a certain percentage of the input on a line-by-line basis:

```
$ seq -f "Line %g" 1000 | sample -r 1%
Line 50
Line 159
Line 173
Line 682
Line 882
Line 921
Line 986
```

Here, every input line has a 1% chance of being printed. This percentage can also be specified as a fraction (namely $1/100$) or as a probability (namely 0.01).

`sample` has two other purposes that can be useful when you're debugging your pipeline. First, it's possible to add some delay to the output. This comes in handy when the input is a constant stream (for example, the Wikipedia stream we saw in [Chapter 3](#)), and the data comes in too fast for you to see what's going on. Second, you can put a timer on `sample` so that you don't have to kill the ongoing process manually. For example, to add a one-second delay between each line being printed and to run for only five seconds, you would type:

```
$ seq -f "Line %g" 1000 | sample -r 1% -d 1000 -s 5 | ts ①
Jun 29 14:28:50 Line 28
Jun 29 14:28:51 Line 262
Jun 29 14:28:52 Line 324
Jun 29 14:28:53 Line 546
Jun 29 14:28:54 Line 589
Jun 29 14:28:55 Line 613
Jun 29 14:28:56 Line 629
```

① The tool `ts`⁹ adds a timestamp in front of each line.

To prevent unnecessary computation, try to put `sample` as early as possible in your pipeline. In fact, this argument holds for any command-line tool that

reduces data, like `head` and `tail`. Once you're confident your pipeline works, you take the tool out of the pipeline.

Extracting Values

To extract the actual chapter headings from our earlier example, you can take a simple approach by piping the output of `grep` to `cut`:¹⁰

```
$ grep -i chapter alice.txt | cut -d ' ' -f 3-
Down the Rabbit-Hole
The Pool of Tears
A Caucus-Race and a Long Tale
The Rabbit Sends in a Little Bill
Advice from a Caterpillar
Pig and Pepper
A Mad Tea-Party
The Queen's Croquet-Ground
The Mock Turtle's Story
The Lobster Quadrille
Who Stole the Tarts?
Alice's Evidence
```

Here, each line that's passed to `cut` is being split on spaces into fields, and then the third field to the last field is being printed. The total number of fields can be different per input line. With `sed` you can accomplish the same task in a much more complex manner:

```
$ sed -rn 's/^CHAPTER ([IVXLCDM]{1,})\. (.*)$/\2/p' alice.txt | trim 3
Down the Rabbit-Hole
The Pool of Tears
A Caucus-Race and a Long Tale
... with 9 more lines
```

This approach uses a regular expression and a back reference. Here, `sed` also takes over the work done by `grep`. I recommend using such a complicated approach only when a simpler one would not work—for example, if *chapter* was ever part of the text itself and not just used to indicate the start of a new chapter. Of course, there are many levels of complexity that would have worked around this, but this is intended to

illustrate an extremely strict approach. In practice, the challenge is to come up with a pipeline that strikes a good balance between complexity and flexibility.

It's worth noting that `cut` can also split on characters' positions. This is useful for when you want to extract (or remove) the same set of characters per input line:

```
$ grep -i chapter alice.txt | cut -c 9-
I. Down the Rabbit-Hole
II. The Pool of Tears
III. A Caucus-Race and a Long Tale
IV. The Rabbit Sends in a Little Bill
V. Advice from a Caterpillar
VI. Pig and Pepper
VII. A Mad Tea-Party
VIII. The Queen's Croquet-Ground
IX. The Mock Turtle's Story
X. The Lobster Quadrille
XI. Who Stole the Tarts?
XII. Alice's Evidence
```

`grep` has a great feature that outputs every match onto a separate line using the `-o` option:

```
$ < alice.txt grep -oE '\w{2,}' | trim
Project
Gutenberg
Alice
Adventures
in
Wonderland
by
Lewis
Carroll
This
... with 28615 more lines
```

But what if you wanted to create a dataset of all the words that start with an *a* and end with an *e*? Well, of course there's a pipeline for that too:

```
$ < alice.txt tr '[:upper:]' '[:lower:]' | ❶
> grep -oE '\w{2,}' |
> grep -E '^a.*e$' |
> sort | uniq | sort -nr | trim
available
ate
assistance
askance
arise
argue
are
archive
applicable
apple
... with 25 more lines
```

- ❶ I use `tr` here to make the text lowercase. We'll have a closer look at `tr` in the next section.

The two `grep` commands might have been combined into one, but in this case I decided it would be easier to reuse and adapt the previous pipeline. There's no shame in being pragmatic to get the job done!

Replacing and Deleting Values

You can use the command-line tool `tr`, which stands for *translate*, to replace or delete individual characters. For example, spaces can be replaced by underscores as follows:

```
$ echo 'hello world!' | tr ' ' '_'
hello_world!
```

If more than one character needs to be replaced, then you can combine those requests:

```
$ echo 'hello world!' | tr ' !' '_?'
hello_world?
```

`tr` can also be used to delete individual characters by specifying the argument `-d`:

```
$ echo 'hello world!' | tr -d ' !'  
helloworld  
  
$ echo 'hello world!' | tr -d -c '[a-z]'  
helloworld%
```

In this case, these two commands accomplish the same thing. The second command, however, uses two additional features: it specifies a *range* of characters (all lowercase letters) using the square brackets and the dash (`[-]`), and the `-c` option indicates that the complement of that should be used. In other words, this command keeps only lowercase letters. You can even use `tr` to convert text to uppercase:

```
$ echo 'hello world!' | tr '[a-z]' '[A-Z]'  
HELLO WORLD!  
  
$ echo 'hello world!' | tr '[:lower:]' '[:upper:]'  
HELLO WORLD!
```

However, if you need to translate non-ASCII characters, then `tr` may not work because it operates on single-byte characters only. In those cases, you should use `sed` instead:

```
$ echo 'hello world!' | tr '[a-z]' '[A-Z]'  
HELLO WORLD!  
  
$ echo 'hallo w reld!' | tr '[a-z]' '[A-Z]'  
HALLO W RELD!  
  
$ echo 'hallo w reld!' | tr '[:lower:]' '[:upper:]'  
HALLO W RELD!  
  
$ echo 'hallo w reld!' | sed 's/[:lower:]*\U&/g'  
HALLO W RELD!  
  
$ echo 'hell  vil g' | tr '[:lower:]' '[:upper:]'  
HELL  VIL G
```

```
$ echo 'helló világ' | sed 's/[:lower:]*$/\U&/g'  
HELLÓ VILÁG
```

If you need to operate on more than individual characters, then you may find `sed` useful. You've already seen an example of `sed` being used to extract the chapter headings from *alice.txt*. Extracting, deleting, and replacing are actually all the same operation in `sed`. You just specify different regular expressions. For example, to change a word, remove repeated spaces, and remove leading spaces:

```
$ echo ' hello      world!' |  
> sed -re 's/hello/bye/' | ❶  
> sed -re 's/\s+/ /g' | ❷  
> sed -re 's/\s+//' ❸  
bye world!
```

- ❶ Replace *hello* with *bye*.
- ❷ Replace any whitespace with one space. The flag `g` stands for *global*, meaning that the same substitution can be applied more than once on the same line.
- ❸ This removes leading spaces only because I didn't specify the flag `g` here.

Again, just as with the `grep` example earlier, these three `sed` commands can be combined into one:

```
$ echo ' hello      world!' |  
> sed -re 's/hello/bye/;s/\s+/ /g;s/\s+//'  
bye world!
```

But tell me—what do you find easier to read?

CSV

The command-line tools such as `tr` and `grep` that I've used to scrub plain text cannot always be applied to CSV. The reason is that these command-line tools have no notion of headers, bodies, and columns.

Bodies and Headers and Columns, Oh My!

What if you want to filter lines using `grep` but always include the header in the output? Or what if you want to uppercase the values of only a specific column using `tr` and leave the other columns untouched?

There are multistep workarounds for this, but they are very cumbersome. I have something better. I'd like to introduce you to three command-line tools, aptly named `body`,¹¹ `header`,¹² and `cols`,¹³ that will enable you to leverage ordinary command-line tools for CSV.

Let's start with the first tool, `body`. With `body` you can apply any command-line tool to the body of a CSV file—that is, to everything excluding the header. For example:

```
$ echo -e "value\n7\n2\n5\n3" | body sort -n
value
2
3
5
7
```

`body` assumes that the header of the CSV file spans only one row. It works like this:

- Take one line from standard input and store it as a variable named `$header`.
- Print out the header.
- Execute all the command-line arguments passed to `body` on the remaining data in standard input.

Here's another example. Imagine that you want to count the lines of the following CSV file:

```
$ seq 5 | header -a count
count
1
2
3
4
5
```

With `wc -l`, you can count the number of all lines:

```
$ seq 5 | header -a count | wc -l
6
```

If you want to consider only the lines in the body (i.e., everything except the header), you add `body`:

```
$ seq 5 | header -a count | body wc -l
count
5
```

Note that the header is not used and is also printed again in the output.

The second command-line tool, `header`, allows you to manipulate the header of a CSV file. If no arguments are provided, the header of the CSV file is printed:

```
$ < tips.csv header
bill,tip,sex,smoker,day,time,size
```

This is the same as `head -n 1`. If the header spans more than one row, which is not recommended, you can specify `-n 2`. You can also add a header to a CSV file:

```
$ seq 5 | header -a count
count
1
2
```

```
3  
4  
5
```

This is equivalent to `echo "count" | cat - <(seq 5)`. Deleting a header is done with the `-d` option:

```
$ < iris.csv header -d | trim  
5.1,3.5,1.4,0.2,Iris-setosa  
4.9,3.0,1.4,0.2,Iris-setosa  
4.7,3.2,1.3,0.2,Iris-setosa  
4.6,3.1,1.5,0.2,Iris-setosa  
5.0,3.6,1.4,0.2,Iris-setosa  
5.4,3.9,1.7,0.4,Iris-setosa  
4.6,3.4,1.4,0.3,Iris-setosa  
5.0,3.4,1.5,0.2,Iris-setosa  
4.4,2.9,1.4,0.2,Iris-setosa  
4.9,3.1,1.5,0.1,Iris-setosa  
... with 140 more lines
```

This is similar to `tail -n +2`, but it's a bit easier to remember. Replacing a header—which basically is deleting a header and then adding a new one if you look at the preceding source code—is accomplished by specifying the `-r` option. Here, we combine it with `body`:

```
$ seq 5 | header -a line | body wc -l | header -r count  
count  
5
```

And last but not least, you can apply a command to just the header, similar to what the `body` command-line tool does to the body. For example:

```
$ seq 5 | header -a line | header -e "tr '[a-z]' '[A-Z]'"  
LINE  
1  
2  
3  
4  
5
```

The third command-line tool is called `cols`, and it allows you to apply a certain command to only a subset of the columns. For example, if you wanted to uppercase the values in the `day` column in the Tips dataset without affecting the other columns and the header, you would use `cols` in combination with `body`, as follows:

```
$ < tips.csv cols -c day body "tr '[a-z]' '[A-Z]'" | head -n 5 | csvlook
  day | bill | tip | sex | smoker | time | size |
  ---+---+---+---+---+---+---+
  0001-01-07 | 16.99 | 1.01 | Female | False | Dinner | 2 |
  0001-01-07 | 10.34 | 1.66 | Male | False | Dinner | 3 |
  0001-01-07 | 21.01 | 3.50 | Male | False | Dinner | 3 |
  0001-01-07 | 23.68 | 3.31 | Male | False | Dinner | 2 |
```

Please note that passing multiple command-line tools and arguments as commands to `header -e`, `body`, and `cols` can lead to tricky quoting citations. If you ever run into such problems, it's best to create a separate command-line tool for this and pass it as command.

In conclusion, while it is generally preferable to use command-line tools that are made specifically for CSV data, `body`, `header`, and `cols` also allow you to apply the classic command-line tools to CSV files if needed.

Performing SQL Queries on CSV

In case the command-line tools mentioned in this chapter do not provide enough flexibility, there is another approach to scrubbing your data from the command line. The tool `csvsql`¹⁴ allows you to execute SQL queries directly on CSV files. SQL is a powerful language for defining operations for scrubbing data; it's a very different approach from using individual command-line tools.

NOTE

If your data originally comes from a relational database, then, if possible, try to execute SQL queries on that database and subsequently extract the data as CSV. As I discussed in [Chapter 3](#), you can use the command-line tool `sql2csv` for this. When you first export data from the database to a CSV file and then apply SQL, not only is it slower, but there is also a possibility that the column types will not be correctly inferred from the CSV data.

In the following scrubbing tasks, I'll include several solutions that involve `csvsql`. A basic command is this:

```
$ seq 5 | header -a val | csvsql --query "SELECT SUM(val) AS sum FROM stdin"
sum
15.0
```

If you pass standard input to `csvsql`, then the table is named `stdin`. The types of the column are automatically inferred from the data. As you'll see later in [“Combining Multiple CSV Files”](#), you can also specify multiple CSV files. Please keep in mind that `csvsql` employs the SQLite dialect of SQL, which has some subtle differences with respect to the SQL standard. While SQL is generally more verbose than the other solutions, it is also much more flexible. If you already know how to tackle a scrubbing problem with SQL, then why not use it when you're at the command line?

Extracting and Reordering Columns

Columns can be extracted and reordered using the command-line tool `csvcut`.¹⁵ For example, to keep only the columns in the Iris dataset that contain numerical values *and* reorder the middle two columns:

```
$ < iris.csv csvcut -c sepal_length,petal_length,sepal_width,petal_width |
  csvlo
ok
| sepal_length | petal_length | sepal_width | petal_width |
|      5.1     |       1.4    |      3.5    |      0.2    |
|      4.9     |       1.4    |      3.0    |      0.2    |
```

	4.7	1.3	3.2	0.2
	4.6	1.5	3.1	0.2
	5.0	1.4	3.6	0.2
	5.4	1.7	3.9	0.4
	4.6	1.4	3.4	0.3
	5.0	1.5	3.4	0.2

... with 142 more lines

Alternatively, you can specify the columns you want to leave out with the `-C` option, which stands for *complement*:

```
$ < iris.csv csvcut -C species | csvlook
+-----+-----+-----+-----+
| sepal_length | sepal_width | petal_length | petal_width |
+-----+-----+-----+-----+
| 5.1 | 3.5 | 1.4 | 0.2 |
| 4.9 | 3.0 | 1.4 | 0.2 |
| 4.7 | 3.2 | 1.3 | 0.2 |
| 4.6 | 3.1 | 1.5 | 0.2 |
| 5.0 | 3.6 | 1.4 | 0.2 |
| 5.4 | 3.9 | 1.7 | 0.4 |
| 4.6 | 3.4 | 1.4 | 0.3 |
| 5.0 | 3.4 | 1.5 | 0.2 |
+-----+-----+-----+-----+
... with 142 more lines
```

Here, the included columns are kept in the same order. Instead of the column names, you can specify the indices of the columns, which start at 1. This allows you to, for example, select only the odd columns (should you ever need to do that!):

```
$ echo 'a,b,c,d,e,f,g,h,i\n1,2,3,4,5,6,7,8,9' |
> csvcut -c $(seq 1 2 9 | paste -sd,)
a,c,e,g,i
1,3,5,7,9
```

If you're certain that there are no commas in any of the values, then you can also use `cut` to extract columns. Be aware that `cut` does not reorder columns, as the following command demonstrates:

```
$ echo 'a,b,c,d,e,f,g,h,i\n1,2,3,4,5,6,7,8,9' | cut -d, -f 5,1,3
a,c,e
1,3,5
```

As you can see, the order in which you specify the columns does not matter with the `-f` option; with `cut`, the columns will always appear in their original order. For completeness, let's also take a look at the SQL approach for extracting and reordering the numerical columns of the Iris dataset:

```
$ < iris.csv csvsql --query "SELECT sepal_length, petal_length, \"\n> \"sepal_width, petal_width FROM stdin" | head -n 5 | csvlook\n| sepal_length | petal_length | sepal_width | petal_width |\n|-----|-----|-----|-----|\n| 5.1 | 1.4 | 3.5 | 0.2 |\n| 4.9 | 1.4 | 3.0 | 0.2 |\n| 4.7 | 1.3 | 3.2 | 0.2 |\n| 4.6 | 1.5 | 3.1 | 0.2 |
```

Filtering Rows

Filtering rows in a CSV file differs from filtering lines in a plain-text file in that you may want to base this filtering only on values in a certain column. Filtering on location is essentially the same, but you have to take into account that the first line of a CSV file is usually the header. Remember that you can always use the `body` command-line tool if you want to keep the header:

```
$ seq 5 | sed -n '3,5p'\n3\n4\n5\n\n$ seq 5 | header -a count | body sed -n '3,5p'\ncount\n3\n4\n5
```

When it comes down to filtering on a certain pattern within a certain column, you can use either `csvgrep`,¹⁶ `awk`, or, of course, `csvsql`. For example, to exclude all the lunch and dinner bills for which the party size was smaller than five people:

```
$ csvgrep -c size -i -r "[1-4]" tips.csv
bill,tip,sex,smoker,day,time,size
29.8,4.2,Female,No,Thur,Lunch,6
34.3,6.7,Male,No,Thur,Lunch,6
41.19,5.0,Male,No,Thur,Lunch,5
27.05,5.0,Female,No,Thur,Lunch,6
29.85,5.14,Female,No,Sun,Dinner,5
48.17,5.0,Male,No,Sun,Dinner,6
20.69,5.0,Male,No,Sun,Dinner,5
30.46,2.0,Male,Yes,Sun,Dinner,5
28.15,3.0,Male,Yes,Sat,Dinner,5
```

Both **awk** and **csvsql** can also do numerical comparisons. For example, to get all the restaurant bills above \$40 on a Saturday or a Sunday:

```
$ < tips.csv awk -F, 'NR==1 || ($1 > 40.0) && ($5 ~ /S/)'
bill,tip,sex,smoker,day,time,size
48.27,6.73,Male,No,Sat,Dinner,4
44.3,2.5,Female,Yes,Sat,Dinner,3
48.17,5.0,Male,No,Sun,Dinner,6
50.81,10.0,Male,Yes,Sat,Dinner,3
45.35,3.5,Male,Yes,Sun,Dinner,3
40.55,3.0,Male,Yes,Sun,Dinner,2
48.33,9.0,Male,No,Sat,Dinner,4
```

The **csvsql** solution is more verbose, but it's also more robust, as it uses the names of the columns instead of their indexes:

```
$ csvsql --query "SELECT * FROM tips WHERE bill > 40 AND day LIKE '%S'"
tips.csv
bill,tip,sex,smoker,day,time,size
48.27,6.73,Male,0,Sat,Dinner,4.0
44.3,2.5,Female,1,Sat,Dinner,3.0
48.17,5.0,Male,0,Sun,Dinner,6.0
50.81,10.0,Male,1,Sat,Dinner,3.0
45.35,3.5,Male,1,Sun,Dinner,3.0
40.55,3.0,Male,1,Sun,Dinner,2.0
48.33,9.0,Male,0,Sat,Dinner,4.0
```

Note that the flexibility of the *WHERE* clause in an SQL query cannot be easily matched with other command-line tools, because SQL can operate on dates and sets and form complex combinations of clauses.

Merging Columns

Merging columns is useful when the values of interest are spread over multiple columns. This may happen with dates (where year, month, and day could be separate columns) or names (where the first name and last name are separate columns). Let's consider the second situation.

The input CSV is a list of composers. Imagine that your task is to combine the first name and the last name into a full name. I'll present four different approaches for this task: `sed`, `awk`, `cols + tr`, and `cvsqsl`. Let's have a look at the input CSV:

```
$ csvlook -I names.csv
+---+---+---+---+
| id | last_name | first_name | born |
+---+---+---+---+
| 1 | Williams | John        | 1932 |
| 2 | Elfman    | Danny       | 1953 |
| 3 | Horner    | James       | 1953 |
| 4 | Shore     | Howard      | 1946 |
| 5 | Zimmer    | Hans        | 1957 |
```

The first approach, `sed`, uses two statements. The first is to replace the header, and the second is a regular expression with back references applied to the second row onward:

```
$ < names.csv sed -re '1s/.*/id,full_name,born/g;2,$s/(.*),(.*),(.*)/\\1,\\3\\2,\\4/g' |
> csvlook -I
+---+---+---+
| id | full_name   | born |
+---+---+---+
| 1 | John Williams | 1932 |
| 2 | Danny Elfman | 1953 |
| 3 | James Horner | 1953 |
| 4 | Howard Shore | 1946 |
| 5 | Hans Zimmer | 1957 |
```

The `awk` approach looks as follows:

```
$ < names.csv awk -F, 'BEGIN{OFS=","; print "id,full_name,born"} {if(NR > 1)
{pr
```

```

int $1,$3" "$2,$4}}' |
> csvlook -I
| id | full_name      | born |
|----|----|----|
| 1  | John Williams  | 1932 |
| 2  | Danny Elfman   | 1953 |
| 3  | James Horner   | 1953 |
| 4  | Howard Shore   | 1946 |
| 5  | Hans Zimmer    | 1957 |

```

Here is the `cols` approach in combination with `tr`:

```

$ < names.csv |
> cols -c first_name,last_name tr \"\",\" \" \" |
> header -r full_name,id,born |
> csvcut -c id,full_name,born |
> csvlook -I
| id | full_name      | born |
|----|----|----|
| 1  | John Williams  | 1932 |
| 2  | Danny Elfman   | 1953 |
| 3  | James Horner   | 1953 |
| 4  | Howard Shore   | 1946 |
| 5  | Hans Zimmer    | 1957 |

```

Please note that `csvsql` employs SQLite as the database to execute the query and that `||` stands for concatenation:

```

$ < names.csv csvsql --query "SELECT id, first_name || ' ' || last_name " \
> "AS full_name, born FROM stdin" | csvlook -I
| id | full_name      | born |
|----|----|----|
| 1.0 | John Williams  | 1932.0 |
| 2.0 | Danny Elfman   | 1953.0 |
| 3.0 | James Horner   | 1953.0 |
| 4.0 | Howard Shore   | 1946.0 |
| 5.0 | Hans Zimmer    | 1957.0 |

```

What if `last_name` could potentially contain a comma? Let's have a look at the raw input CSV for clarity's sake:

```

$ cat names-comma.csv
id,last_name,first_name,born

```

```
1,Williams,John,1932
2,Elfman,Danny,1953
3,Horner,James,1953
4,Shore,Howard,1946
5,Zimmer,Hans,1957
6,"Beethoven, van",Ludwig,1770
```

Well, it appears that the first three approaches fail, all in different ways. Only `csvsql` is able to combine `first_name` and `full_name`:

```
$ < names-comma.csv sed -re '1s/.*/id,full_name,born/g;2,$s/(.*),(.*),(.*)/\\1,\\3 \\2,\\4/g' | tail -n 1
6,"Beethoven,Ludwig van",1770

$ < names-comma.csv awk -F, 'BEGIN{OFS=","; print "id,full_name,born"} {if(NR >
1) {print $1,$3" "$2,$4}}' | tail -n 1
6, van" Beethoven,Ludwig

$ < names-comma.csv cols -c first_name,last_name tr '\',\' \' \' |
> header -r full_name,id,born | csvcut -c id,full_name,born | tail -n 1
6,"Ludwig ""Beethoven van""",1770

$ < names-comma.csv csvsql --query "SELECT id, first_name || ' ' || last_name
AS
full_name, born FROM stdin" | tail -n 1
6.0,"Ludwig Beethoven, van",1770.0

$ < names-comma.csv rush run -t 'unite(df, full_name, first_name, last_name,
sep
= " ")' - | tail -n 1
6,"Ludwig Beethoven, van",1770
```

Wait a minute! What's that last command? Is that R? Well, as a matter of fact, it is. It's R code evaluated through the command-line tool `rush`. All that I can say at this moment is that this approach also succeeds at merging the two columns. I'll discuss this nifty command-line tool later.

Combining Multiple CSV Files

If you start out with multiple CSV files, it's often a good idea to combine them. When the CSV files have the same columns, you can concatenate them horizontally. When the CSV files have different column names but are still related, you can often join them.

Concatenate horizontally

Let's say you have three CSV files that you want to put side by side. We use `tee`¹⁷ to save the result of `csvcut` in the middle of the pipeline:

```
$ < tips.csv csvcut -c bill,tip | tee bills.csv | head -n 3 | csvlook
| bill | tip |
|-----|
| 16.99 | 1.01 |
| 10.34 | 1.66 |

$ < tips.csv csvcut -c day,time | tee datetime.csv |
> head -n 3 | csvlook -I
| day | time |
|-----|
| Sun | Dinner |
| Sun | Dinner |

$ < tips.csv csvcut -c sex,smoker,size | tee customers.csv |
> head -n 3 | csvlook
| sex | smoker | size |
|-----|
| Female | False | 2 |
| Male | False | 3 |
```

Assuming that the rows line up, you can `paste` the files together:

```
$ paste -d, {bills,customers,datetime}.csv | head -n 3 | csvlook -I
| bill | tip | sex | smoker | size | day | time |
|-----|
| 16.99 | 1.01 | Female | No | 2 | Sun | Dinner |
| 10.34 | 1.66 | Male | No | 3 | Sun | Dinner |
```

Here the command-line argument `-d` instructs `paste` to use a comma as the delimiter.

Joining

Sometimes data cannot be combined by vertical or horizontal concatenation. In some cases, especially in relational databases, the data is spread over multiple tables (or files) to minimize redundancy. Imagine you wanted to extend the Iris dataset with more information about the three types of iris flowers, namely the USDA identifiers. It so happens that I have a separate CSV file with these identifiers:

```
$ csvlook irismeta.csv
| species | wikipedia_url | usda_id |
| Iris-versicolor | http://en.wikipedia.org/wiki/Iris_versicolor | IRVE2 |
| Iris-virginica | http://en.wikipedia.org/wiki/Iris_virginica | IRVI |
| Iris-setosa | | IRSE |
```

What this dataset and the Iris dataset have in common is the *species* column. You can use `csvjoin`¹⁸ to join the two datasets:

```
$ csvjoin -c species iris.csv irismeta.csv | csvcut -c
sepal_length,sepal_width,
species,usda_id | sed -n '1p;49,54p' | csvlook
| sepal_length | sepal_width | species | usda_id |
| 4.6 | 3.2 | Iris-setosa | IRSE |
| 5.3 | 3.7 | Iris-setosa | IRSE |
| 5.0 | 3.3 | Iris-setosa | IRSE |
| 7.0 | 3.2 | Iris-versicolor | IRVE2 |
| 6.4 | 3.2 | Iris-versicolor | IRVE2 |
| 6.9 | 3.1 | Iris-versicolor | IRVE2 |
```

Of course, you can also opt for the SQL approach using `csvsql`, which is, as per usual, a bit longer (but potentially much more flexible):

```
$ csvsql --query 'SELECT i.sepal_length, i.sepal_width, i.species, m.usda_id
FROM
iris i JOIN irismeta m ON (i.species = m.species)' iris.csv irismeta.csv |
sed
-n '1p;49,54p' | csvlook
| sepal_length | sepal_width | species | usda_id |
| 4.6 | 3.2 | Iris-setosa | IRSE |
```

5.3	3.7	Iris-setosa	IRSE
5.0	3.3	Iris-setosa	IRSE
7.0	3.2	Iris-versicolor	IRVE2
6.4	3.2	Iris-versicolor	IRVE2
6.9	3.1	Iris-versicolor	IRVE2

Working with XML/HTML and JSON

In this section, I'm going to demonstrate a couple of command-line tools that can convert data from one format to another. There are two reasons to convert data.

First, oftentimes the data needs to be in tabular form, just like a database table or a spreadsheet, because many visualization and machine learning algorithms depend on it. CSV is inherently in tabular form, but JSON and HTML/XML data can have a deeply nested structure.

Second, many command-line tools, especially the classic ones such as `cut` and `grep`, operate on plain text. This is because text is regarded as a universal interface between command-line tools. Moreover, the other formats are younger. Each of these formats can be treated as plain text, allowing us to apply such command-line tools to the other formats as well.

Sometimes you can get away with applying the classic tools to structured data. For example, by treating the following JSON data as plain text, you can change the attribute `gender` to `sex` using `sed`:

```
$ sed -e 's/"gender":/"sex":/g' users.json | jq | trim
{
  "results": [
    {
      "sex": "male",
      "name": {
        "title": "mr",
        "first": "leevi",
        "last": "kivisto"
      },
      "location": {
... with 260 more lines
```

Like many other command-line tools, `sed` does not make use of the structure of the data. Better to either use a tool that makes use of the structure of the data (such as `jq`, which I'll discuss in a moment) or to first convert the data to a tabular format such as CSV and then apply the appropriate command-line tool.

I'm going to demonstrate converting XML/HTML and JSON to CSV through a real-world use case. The command-line tools that I'll be using here are `curl`, `pup`, `xml2json`,¹⁹ `jq`, and `json2csv`.²⁰

Wikipedia holds a wealth of information. Much of this information is ordered in tables, which can be regarded as datasets. For example, [this page](#) contains a list of countries and territories together with their border lengths and surface areas and the ratio between the two.

Let's imagine that you're interested in analyzing this data. In this section, I'll walk you through all the necessary steps and their corresponding commands. I won't go into every little detail, so it could be that you won't understand everything right away. Don't worry—I'm confident that you'll get the gist of it. Remember that the purpose of this section is to demonstrate the command line. All tools and concepts used in this section (and more) will be explained in the subsequent chapters.

The dataset that you're interested in is embedded in HTML. Your goal is to end up with a representation of this dataset that you can work with. The very first step is to download the HTML using `curl`:

```
$ curl -sL  
'http://en.wikipedia.org/wiki/List_of_countries_and_territories_by_bo  
rder/area_ratio' > wiki.html
```

The HTML is saved to a file named `wiki.html`. Let's see what the first 10 lines look like:

```
$ < wiki.html trim  
<!DOCTYPE html>  
<html class="client-nojs" lang="en" dir="ltr">  
<head>  
<meta charset="UTF-8"/>
```

```
<title>List of countries and territories by border/area ratio -  
Wikipedia</titl...  
<script>document.documentElement.className="client-js";RLCONF=  
{"wgBreakFrames":...  
"Lists of countries by geography","Lists by area","Border-related  
lists"], "wgPa...  
!1,"wgGELinkRecommendationsFrontendEnabled":!1,"wgWikibaseItemId": "Q6613807"};  
R...  
"ext.navigationTiming","ext.uls.compactlinks","ext.uls.interface","ext.cx.even  
t...  
<script>(RLQ=window.RLQ||[]).push(function()  
{mw.loader.implement("user.options@...  
... with 3038 more lines
```

That seems to be in order. Now image that you've been able to determine that the root HTML element that you're interested in is a `<table>` with the class `wikitable`. This allows you to look at the part you're interested in using `grep` (the `-A` option specifies the number of lines you want to print after the matching line):

```
$ grep wikitable -A 21 wiki.html  
<table class="wikitable sortable">  
<tbody><tr>  
<th>Rank</th>  
<th>Country or territory</th>  
<th>Total length of land borders (km)</th>  
<th>Total surface area (km2)</th>  
<th>Border/area ratio (km/km2)</th>  
</th></tr>  
<tr>  
<td>1  
</td>  
<td>Vatican City  
</td>  
<td>3.2  
</td>  
<td>0.44  
</td>  
<td>7.2727273  
</td></tr>  
<tr>  
<td>2  
</td>
```

You now actually see the countries and their values. The next step is to extract the necessary elements from the HTML file. For this you can use `pup`:

```
$ < wiki.html pup 'table.wikitable tbody' | tee table.html | trim
<tbody>
<tr>
<th>
  Rank
</th>
<th>
  Country or territory
</th>
<th>
  Total length of land borders (km)
... with 4199 more lines
```

The expression passed to `pup` is a CSS selector. The syntax is usually used to style web pages, but you can also use it to select certain elements from HTML. In this case, you want to select the `tbody` of the `table` that has the `wikitable` class. Up next is `xml2json`, which converts XML (and HTML) to JSON:

```
$ < table.html xml2json > table.json

$ jq -c table.json | trim 20
{
  "tbody": {
    "tr": [
      {
        "th": [
          {
            "$t": "Rank"
          },
          {
            "$t": "Country or territory"
          },
          {
            "$t": "Total length of land borders (km)"
          }
        ],
        "td": [
          "Total surface area (km",
          "Population (2016 est.)",
          "GDP (nominal) (2016 est.)"
        ]
      }
    ]
  }
}
```

```
        ")"
],
"sup": {
... with 4691 more lines
```

The reason you convert the HTML to JSON is because there is a very powerful tool called `jq` that operates on JSON data. The following command extracts certain parts of the JSON data and reshapes it into a form that you can work with:

```
$ < table.json jq -r '.tbody.tr[1:][] | [.td[]["$t"]]' | @csv' | header -a
rank,c
ountry,border,surface,ratio > countries.csv
```

The data is now in a form that you can work with. Those were quite a few steps to get from a Wikipedia page to a CSV dataset. However, when you combine all of the preceding commands into one, you see that it's actually really concise and expressive:

```
$ csvlook --max-column-width 28 countries.csv
+-----+-----+-----+-----+-----+
| rank | country | border | surface | ratio |
+-----+-----+-----+-----+-----+
| 1 | Vatican City | 3.20 | 0.44 | 7.273... |
| 2 | Monaco | 4.40 | 2.00 | 2.200... |
| 3 | San Marino | 39.00 | 61.00 | 0.639... |
| 4 | Liechtenstein | 76.00 | 160.00 | 0.465... |
| 5 | Sint Maarten (Netherlands) | 10.20 | 34.00 | 0.300... |
| 6 | Andorra | 120.30 | 468.00 | 0.257... |
| 7 | Gibraltar (United Kingdom) | 1.20 | 6.00 | 0.200... |
| 8 | Saint Martin (France) | 10.20 | 54.00 | 0.189... |
+-----+-----+-----+-----+-----+
... with 238 more lines
```

That concludes the demonstration of converting XML/HTML to JSON to CSV. While `jq` can perform many more operations, and while there exist specialized tools to work with XML data, in my experience, converting the data to CSV format as quickly as possible tends to work well. This way, you can spend more time becoming proficient at using generic command-line tools rather than very specific tools.

Summary

In this chapter we've looked at cleaning or scrubbing data. As you've seen, there is no single tool that can magically get rid of all the messiness of data; you'll often need to combine different tools to get the desired result. Keep in mind that classic command-line tools such as `cut` and `sort` can't interpret structured data. Luckily, there are tools that convert one data format, such as JSON and XML, into another data format, such as CSV. In the next chapter, which is again an intermezzo chapter, I'm going to show you how you can manage your project using `make`. You're free to skip Chapter 6 if you can't wait to start exploring and visualizing your data in Chapter 7.

For Further Exploration

- I wish I could've explained more about `awk`. It's such a powerful tool and programming language. I highly recommend that you take the time to learn it. Two good resources for that are the book *sed & awk* by Dale Doherty and Arnold Robbins (O'Reilly) and the online [GNU Awk User's Guide](#).
- In this chapter I have used regular expressions in a couple of places. A tutorial about them is unfortunately beyond the scope of this book. Because regular expressions can be used in many different tools, I recommend that you learn about them. A good book for this is *Regular Expressions Cookbook* by Jan Goyvaerts and Steven Levithan (O'Reilly).

¹ Mike D. Brennan and Thomas E. Dickey, *awk – Pattern Scanning and Text Processing Language*, version 1.3.4, 2019, <https://invisible-island.net/mawk>.

² Jeroen Janssens, *rush – R One-Liners from the Shell*, version 0.1, 2021, <https://github.com/jeroenjanssens/rush>.

³ The Linux Information Project, “Plain Text Definition,” last updated February 9, 2007, http://www.linfo.org/plain_text.html.

- 4 Andrew Hunt and David Thomas, *The Pragmatic Programmer* (Reading, MA: Addison-Wesley, 1999).
- 5 Raymond, *The Art of Unix Programming*.
- 6 Jay Fenlason et al., *sed – Stream Editor for Filtering and Transforming Text*, version 4.7, 2018, <https://www.gnu.org/software/sed>.
- 7 Paul Rubin et al., *tail – Output the Last Part of Files*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 8 Jeroen Janssens, *sample – Filter Lines from Standard Input According to Some Probability, with a Given Delay, and for a Certain Duration*, version 0.2.4, 2021, <https://github.com/jeroenjanssens/sample>.
- 9 Joey Hess, *ts – Timestamp Input*, version 0.65, 2021, <https://joeyh.name/code/moreutils>.
- 10 David M. Ihnat, David MacKenzie, and Jim Meyering, *cut – Remove Sections from Each Line of Files*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 11 Jeroen Janssens, *body – Apply Command to All but the First Line*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.
- 12 Jeroen Janssens, *header – Add, Replace, and Delete Header Lines*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.
- 13 Jeroen Janssens, *cols – Apply Command to Subset of Columns*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.
- 14 Christopher Groskopf, *csvsql – Execute SQL Statements on CSV Files*, version 1.0.5, 2020, <https://csvkit.rtfd.org>.
- 15 Christopher Groskopf, *csvcut – Filter and Truncate CSV Files*, version 1.0.5, 2020, <https://csvkit.rtfd.org>.
- 16 Christopher Groskopf, *csvgrep – Search CSV Files*, version 1.0.5, 2020, <https://csvkit.rtfd.org>.
- 17 Mike Parker, Richard M. Stallman, and David MacKenzie, *tee – Read from Standard Input and Write to Standard Output and Files*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 18 Christopher Groskopf, *csvjoin – Execute a SQL-Like Join to Merge CSV Files on a Specified Column or Columns*, version 1.0.5, 2020, <https://csvkit.rtfd.org>.
- 19 François Parmentier, *xml2json – Convert an XML Input to a JSON Output, Using xml-mapping*, version 0.0.3, 2016, <https://github.com/parmentf/xml2json>.
- 20 Jehiah Czebotar, *json2csv – Convert JSON to CSV*, version 1.2.1, 2019, <https://github.com/jehiah/json2csv>.

Chapter 6. Project Management with Make

I hope that by now you have come to appreciate that the command line is a very convenient environment for working with data. You may have noticed that, as a consequence of working with the command line, we:

- Invoke many different commands
- Work from various directories
- Develop our own command-line tools
- Obtain and generate many (intermediate) files

Since this is an exploratory process, our workflow tends to be rather chaotic, which makes it difficult to keep track of what we've done. It's important that our steps can be reproduced, both by us and by others. When you continue with a project from some time ago, chances are that you have forgotten which commands you ran, from which directory, on which files, with which parameters, and in which order. Imagine the challenges of sharing your project with a collaborator.

You can recover some commands by digging through the output of the `history` command, but this is, of course, not a reliable approach. A somewhat better approach would be to save your commands to a shell script. At least this allows you and your collaborators to reproduce the project. A shell script is, however, also suboptimal, for several reasons:

- It is difficult to read and to maintain.
- Dependencies between steps are unclear.
- Every step gets executed every time, which is inefficient and is also sometimes undesirable.

This is where `make` really shines. `make`¹ is a command-line tool that allows you to:

- Formalize your data workflow steps in terms of input and output dependencies
- Run specific steps of your workflow
- Use inline code
- Store and retrieve data from external sources

NOTE

In the first edition, this chapter used `dmake`² instead of `make`. `dmake` was supposed to be a successor to `make` with additional features for working with data. Unfortunately, `dmake` was abandoned by its creators in 2016 with too many unresolved bugs. That's why I've decided to use `make` instead.

An important and related topic is *version control*, which allows you to track changes to your project, back up your project to a server, collaborate with others, and retrieve earlier versions of your project when things go wrong. A popular command-line tool for doing version control is `git`. It's often used in combination with GitHub, an online service for distributed version control. Many open source projects, including [this book](#), are hosted on GitHub. The topic of version control is beyond the scope of this book, but I highly recommend that you look into this, especially once you start collaborating with others. At the end of this chapter I recommend a few resources for learning more.

Overview

Managing your data workflow with `make` is the main topic of this chapter. As such, you'll learn about:

- Defining your workflow with a *Makefile*

- Thinking about your workflow in terms of input and output dependencies
- Running tasks and building targets

This chapter starts with the following files:

```
$ cd /data/ch06
$ l
total 28K
-rw-r--r-- 1 dst dst  37 Jun 29 14:29 Makefile.test
-rw-r--r-- 1 dst dst  16 Jun 29 14:29 numbers.make
-rw-r--r-- 1 dst dst  26 Jun 29 14:29 numbers-write.make
-rw-r--r-- 1 dst dst  21 Jun 29 14:29 numbers-write-var.make
-rw-r--r-- 1 dst dst 432 Jun 29 14:29 starwars.make
-rw-r--r-- 1 dst dst 263 Jun 29 14:29 tasks.make
-rw-r--r-- 1 dst dst  27 Jun 29 14:29 template.make
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Introducing Make

`make` organizes command execution around data and its dependencies. Your data processing steps are formalized in a separate text file (a workflow). Each step may have inputs and outputs. `make` automatically resolves their dependencies and determines which commands need to be run and in which order.

This means that when you have, say, an SQL query that takes 10 minutes, it only has to be executed when the result is missing or when the query has changed afterward. Also, if you want to (re)run a specific step, `make` reruns only the steps on which that step depends. This can save you a lot of time.

Having a formalized workflow allows you to easily pick up your project after a few weeks and to collaborate with others. I strongly advise you to have one, even when you think something will be a one-off project, because

you never know when you might need to run certain steps again or reuse them in another project.

Running Tasks

By default, `make` searches for a configuration file called *Makefile* in the current directory. It can also be named *makefile* (lowercase), but I recommend calling your file *Makefile* because it's more common and because it appears at the top of a directory listing that way. Normally, you would have only one configuration file per project. Because this chapter discusses many different ones, I have given each of them a different filename with the *.make* extension. Let's start with the following *Makefile*:

```
$ bat -A numbers.make
-----
| File: numbers.make
-----
| 1 | numbers: L_F
| 2 |   seq 7 L_F
-----
```

This *Makefile* contains one *target* called `numbers`. A target is like a task. It's usually the name of a file you'd like to create, but it can also be more generic than that. The line below, `seq 7`, is known as a *rule*. Think of a rule as a recipe; it consists of one or more commands that specify how the target should be built.

The whitespace in front of the rule is a single tab character. `make` is picky when it comes to whitespace. Be aware that some editors insert spaces when you press the Tab key, known as a “soft tab,” which will cause `make` to produce an error. The following code illustrates this by expanding the tab to eight spaces:

```
$ < numbers.make expand > spaces.make
```

```
$ bat -A spaces.make
-----
| File: spaces.make
-----
1 | numbers: L_F
2 | .....seq·7 L_F
-----
$ make -f spaces.make ①
spaces.make:2: *** missing separator (did you mean TAB instead of 8 spaces?).
S
top. ②

$ rm spaces.make
```

- ① I need to add the `-f` option (short for the `--makefile` option) because the configuration file isn't called *Makefile*, which is the default.
- ② One of the more helpful error messages you'll find at the command line!

From now on, I'll rename the appropriate file *Makefile* because that matches real-world use more closely. So if I just run `make`:

```
$ cp numbers.make Makefile
$ make
seq 7
1
2
3
4
5
6
7
```

then we see that `make` first prints the rule itself (`seq 7`) and then prints the output generated by the rule. This process is known as *building* a target. If

you don't specify the name of a target, then `make` will build the first target specified in the *Makefile*. In practice, though, you'll most often specify the target you want to build:

```
$ make numbers
seq 7
1
2
3
4
5
6
7
```

NOTE

`make` was originally created to ease the compilation of source code, which explains some of the terminology such as *target*, *rule*, and *building*.

In this case, we're not actually building anything—that is, we're not creating any new files. `make` will happily *build* our target `numbers` again, because it's not finding a file called `numbers`. I'll go into this in the next section.

Sometimes it's useful to have a target that builds regardless of whether a file with the same name exists. Think of tasks that you need to perform as part of a project. It's good practice to declare those targets as phony by using a special target called `.PHONY` at the top of your *Makefile*, followed by the names of the phony targets. Here's an example *Makefile* that illustrates the use of phony targets:

```
$ bat tasks.make
-----
| File: tasks.make
-----
1 | .PHONY: clean publish docker-run
2 |
```

```
3 | clean:
4 |     rm book/2e/book.md book/2e/render*.rds
5 |
6 | publish:
7 |     (cd www && hugo) && netlify deploy --prod --dir www/public
8 |
9 | docker-run:
10 |    docker run -it --rm -v $$(pwd)/book/2e/data:/data -p
11 |    8000:8000
12 |    datasciencetoolbox/dsatcl2e:latest ①
```

- ① Note the extra dollar sign in front of `$(pwd)`. This is needed because `make` uses a single dollar sign to refer to various special variables, which I'll explain later.

This is taken from a *Makefile* I've used while working on this book. You could say that I'm using `make` as a glorified task runner. Although this isn't the primary purpose of `make`, it still provides a lot of value, because I don't need to remember or look up what incantation I used. Instead, I type `make publish`, and the latest version of the book is published. It's perfectly fine to put long-running commands in a *Makefile*.

And `make` can do much more for us!

Building, for Real

Let's modify our *Makefile* so that the output of the rule is written to a file called *numbers*:

```
$ cp numbers-write.make Makefile
$ bat Makefile
-----
| File: Makefile
-----
```

```

1 | numbers:
2 |     seq 7 > numbers


---


—
$ make numbers
seq 7 > numbers

$ bat numbers


---


—
| File: numbers


---


—
1 | 1
2 | 2
3 | 3
4 | 4
5 | 5
6 | 6
7 | 7


---


—

```

Now we can say that `make` is actually building something. What's more, if we run it again, we see that `make` reports that the target `numbers` is up-to-date:

```
$ make numbers
make: 'numbers' is up to date.
```

There's no need to rebuild the target `numbers` because the file *numbers* already exists. That's great, because `make` is saving us time by not repeating work.

In `make`, it's all about files. But keep in mind that `make` only cares about the *name* of the target. It does not check whether a file of the same name actually gets created by the rule. If we were to write to a file called *numbers*, which is Dutch for “numbers,” and the target was still called `numbers`, then `make` would always build this target. Vice versa, if the file `numbers` was created by some other process, whether automated or manual, then `make` would still consider that target up-to-date.

We can avoid some repetition by using the automatic variable `$@`, which gets expanded to the name of the target:

```
$ cp numbers-write-var.make Makefile
```

```
$ bat Makefile
```

```
-----  
|  
|----- File: Makefile  
|  
|-----  
| 1 | numbers:  
| 2 |     seq 7 > $@  
|-----  
|  
-----
```

Let's verify that this works by removing the file *numbers* and calling `make` again:

```
$ rm numbers
```

```
$ make numbers  
seq 7 > numbers
```

```
$ bat numbers
```

```
-----  
|  
|----- File: numbers  
|  
|-----  
| 1 | 1  
| 2 | 2  
| 3 | 3  
| 4 | 4  
| 5 | 5  
| 6 | 6  
| 7 | 7  
|-----  
|  
-----
```

Another reason for `make` to rebuild a target is its dependencies, so let's discuss that next.

Adding Dependencies

So far, we've looked at targets that exist in isolation. In a typical data science workflow, many steps depend on other steps. In order to properly talk about dependencies in a *Makefile*, let's consider two tasks that work with a dataset about *Star Wars* characters.

Here's an excerpt from that dataset:

```
$ curl -sL 'https://raw.githubusercontent.com/tidyverse/dplyr/master/data-
raw/st
arwars.csv' |
> xsv select name,height,mass,homeworld,species |
> csvlook
| name           | height | mass   | homeworld | species |
|:---|:---|:---|:---|:---|
| Luke Skywalker | 172    | 77.0  | Tatooine  | Human   |
| C-3PO          | 167    | 75.0  | Tatooine  | Droid    |
| R2-D2          | 96     | 32.0  | Naboo     | Droid    |
| Darth Vader    | 202    | 136.0 | Tatooine  | Human   |
| Leia Organa    | 150    | 49.0  | Alderaan  | Human   |
| Owen Lars      | 178    | 120.0 | Tatooine  | Human   |
| Beru Whitesun lars | 165    | 75.0  | Tatooine  | Human   |
| R5-D4          | 97     | 32.0  | Tatooine  | Droid    |
... with 79 more lines
```

The first task computes the 10 tallest humans:

```
$ curl -sL 'https://raw.githubusercontent.com/tidyverse/dplyr/master/data-
raw/st
arwars.csv' |
> grep Human | ①
> cut -d, -f 1,2 | ②
> sort -t, -k2 -nr | ③
> head ④
Darth Vader,202
Qui-Gon Jinn,193
Dooku,193
Bail Prestor Organa,191
Raymus Antilles,188
Mace Windu,188
Anakin Skywalker,188
Gregar Typho,185
```

```
Jango Fett,183  
Clegg Lars,183
```

- ❶ Only keep lines that contain the pattern Human.
- ❷ Extract the first two columns.
- ❸ Sort the lines by the second column in reverse numeric order.
- ❹ By default, `head` prints the first 10 lines. You can override this with the `-n` option.

The second task creates a box plot showing the distribution of heights per species (see [Figure 6-1](#)):

```
$ curl -sL 'https://raw.githubusercontent.com/tidyverse/dplyr/master/data-  
raw/st  
arwars.csv' |  
> rush plot --x height --y species --geom boxplot > heights.png  
  
$ display heights.png
```

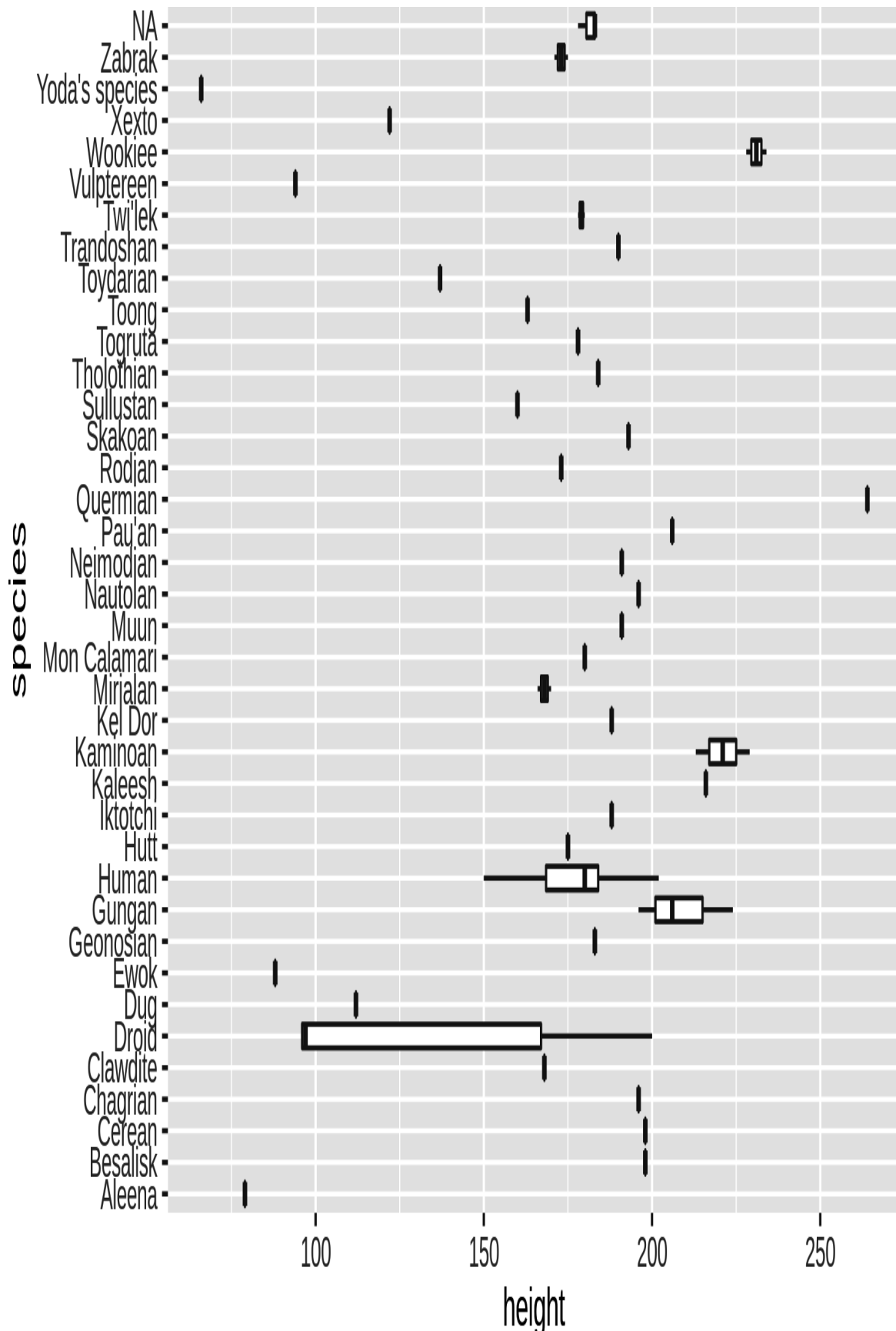


Figure 6-1. Distribution of heights per species in Star Wars

Let's put these two tasks into a *Makefile*. Instead of doing this incrementally, I'd first like to show what a complete *Makefile* looks like and then explain all of the syntax step by step:

```
$ cp starwars.make Makefile
$ bat Makefile
-----
| File: Makefile
-----
|
1 | SHELL := bash
2 | .ONESHELL:
3 | .SHELLFLAGS := -eu -o pipefail -c
4 |
5 | URL = "https://raw.githubusercontent.com/tidyverse/dplyr/master/data-
ra
   | w/starwars.csv"
6 |
7 | .PHONY: all top10
8 |
9 | all: top10 heights.png
10 |
11 | data:
12 |     mkdir $@
13 |
14 | data/starwars.csv: data
15 |     curl -sL $(URL) > $@
16 |
17 | top10: data/starwars.csv
18 |     grep Human $< |
19 |     cut -d, -f 1,2 |
20 |     sort -t, -k2 -nr |
21 |     head
22 |
23 | heights.png: data/starwars.csv
24 |     < $< r ush plot --x height --y species --geom boxplot > $@
```

Now let's go through this *Makefile* step by step. The first three lines are there to change some default settings related to `make` itself:

1. All rules are executed in a shell, which, by default, is `sh`. With the `SHELL` variable, we can change this to another shell, like `bash`. This way, we can use everything that Bash has to offer, such as `for` loops.
2. By default, every line in a rule is sent separately to the shell. With the special target `.ONESHELL`, we can override this so that the rule for the target `top10` works.
3. The `.SHELLFLAGS` line makes Bash more strict, which is considered a **best practice**. For example, because of this, the pipeline in the rule for the target `top10` now stops as soon as there is an error.

We define a custom variable called `URL`. Even though this is used only once, I find it helpful to put information like this near the beginning of the file so that you can easily make changes to these kinds of settings.

With the special target `.PHONY`, we can indicate which targets are not represented by files. In our case, that holds true for the targets `all` and `top10`. These targets will now be executed regardless of whether the directory contains files with the same name.

There are five targets: `all`, `data`, `data/starwars.csv`, `top10`, and `heights.png`. **Figure 6-2** provides an overview of these targets and the dependencies between them.

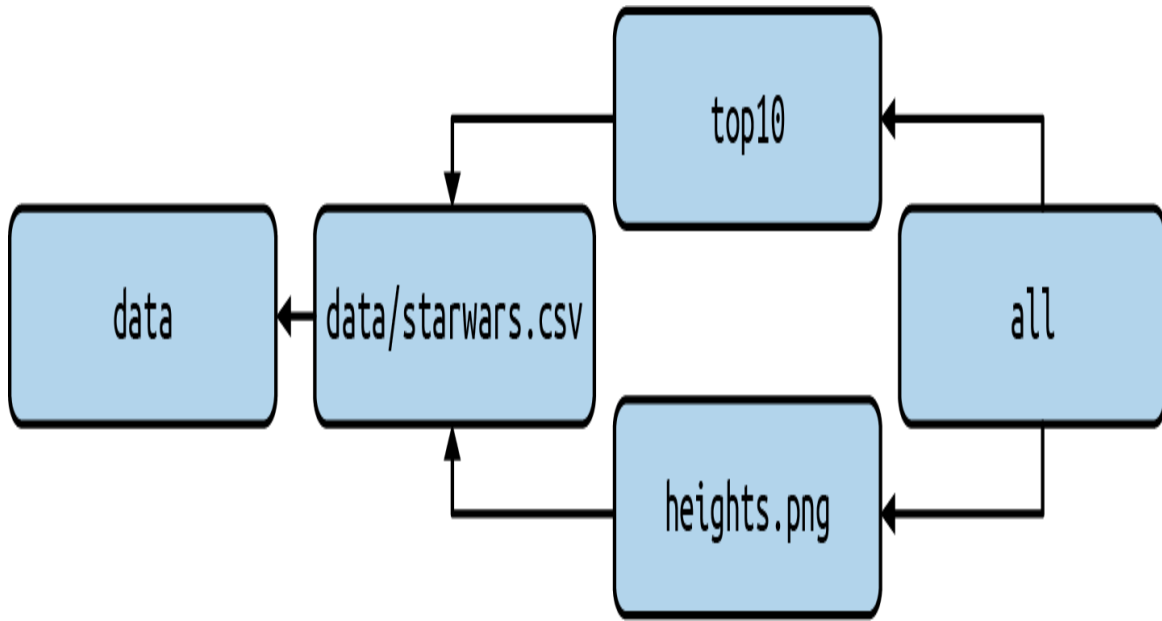


Figure 6-2. Dependencies between targets

Let's discuss each target in turn:

1. The target `all` has two dependencies but no rule. This is like a shortcut to execute one or more targets in the order in which they are specified—in this case, `top10` and `heights.png`. The target `all` appears as the first target in the *Makefile*, which means that if we run `make`, this target will be built.
2. The target `data` creates the directory `data`. Earlier I said that `make` is all about files. Well, it's also about directories. This target will be executed only if the directory `data` doesn't yet exist.
3. The target `data/starwars.csv` depends on the target `data`. If there's no `data` directory, it will first be created. Once all dependencies are satisfied, the rule will be executed, which involves downloading a file and saving it to a file with the same name as the target.
4. The target `top10` is marked as phony, so it will always be built if specified. It depends on the `data/starwars.csv` target. It makes use of a special variable, `$<`, which expands to the name of the first prerequisite, namely `data/starwars.csv`.

5. The target `heights.png`, like the target `top10`, depends on `data/starwars.csv` and makes use of both automatic variables we've seen in this chapter. See the [online documentation](#) if you'd like to learn about other automatic variables.

Last but not least, let's verify that this *Makefile* works:

```
$ make
mkdir data
curl -sL "https://raw.githubusercontent.com/tidyverse/dplyr/master/data-
raw/star-
wars.csv" > data/starwars.csv
grep Human data/starwars.csv |
cut -d, -f 1,2 |
sort -t, -k2 -nr |
head
Darth Vader,202
Qui-Gon Jinn,193
Dooku,193
Bail Prestor Organa,191
Raymus Antilles,188
Mace Windu,188
Anakin Skywalker,188
Gregar Typho,185
Jango Fett,183
Cliegg Lars,183
< data/starwars.csv rush plot --x height --y species --geom boxplot >
heights.png
g
```

No surprises here. Because we didn't specify any target, the `all` target will be built, which in turn causes both the `top10` and `heights.png` targets to be built. The output of the former is printed to standard output, and the latter creates the file `heights.png`. The `data` directory is created only once, just like the CSV file is downloaded only once.

There's nothing more fun than just playing with your data and forgetting everything else. But you have to trust me when I say that it's worthwhile to keep a record of what you have done using a *Makefile*. Not only will it make your life easier (pun intended), but you will also start thinking about your data workflow in terms of steps. Just as with your own command-line

toolbox, which you expand over time, the same holds for `make` workflows. The more steps you have defined, the easier it gets to keep doing it, because very often you can reuse certain steps. I hope that you will get used to `make`, and that it will make your life easier.

Summary

One of the beauties of the command line is that it allows you to play with your data. You can easily execute different commands and process different datafiles. It is a very interactive and iterative process. After a while, it is easy to forget which steps you have taken to get the desired result. It's therefore very important to document your steps every once in a while. This way, if you or one of your colleagues picks up your project after some time, the same result can be produced again by executing the same steps.

In this chapter I've shown you that just putting every command in one Bash script is suboptimal. Instead, I propose that you use `make` as a command-line tool to manage your data workflow. The next chapter covers the third step of the OSEMN model for data science: exploring data.

For Further Exploration

- The book *Managing Projects with GNU Make* by Robert Mecklenburg (O'Reilly) and the online *GNU Make Manual* provide a comprehensive and advanced overview of `make`.
- There are plenty of other workflow managers besides `make`. Although they differ in syntax and features, they all use concepts such as targets, rules, and dependencies. Examples include [Luigi](#), [Apache Airflow](#), and [Nextflow](#).
- To learn more about version control, and about `git` and GitHub in particular, I recommend the book *Pro Git* by Scott Chacon and Ben Straub (Apress); it's [available for free](#). The [online GitHub documentation](#) is also a great starting point.

1 Stuart I. Feldman, *make – A Program for Maintaining Computer Programs*, version 4.3, 2020,
<https://www.gnu.org/software/make>.

2 Factual, *drake – Data Workflow Tool, like a “Make for Data,”* version 1.0.3, 2016,
<https://github.com/Factual/drake>.

Chapter 7. Exploring Data

After all that hard work (unless you already had clean data lying around), it's time for some fun. Now that you have obtained and scrubbed your data, you can continue with the third step of the OSEMN model, which is to explore your data.

Exploring is the step where you familiarize yourself with the data. Being familiar with the data is essential when you want to extract any value from it. For example, knowing what kind of features the data has means you know which features are worth further exploration and which ones you can use to answer any questions that you have.

Exploring your data can be done from three perspectives. The first perspective is to inspect the data and its properties. Here, you want to find out things like what the raw data looks like, how many data points the dataset has, and which features the dataset has.

The second is to compute descriptive statistics. This perspective is useful for learning more about the individual features. The output is often brief and textual and can therefore be printed on the command line.

The third perspective is to create visualizations of the data. From this perspective you can gain insight into how multiple features interact. I'll discuss a way of creating visualizations that can be printed on the command line. However, visualizations are best suited for display on a GUI. An advantage of data visualizations over descriptive statistics is that the former are more flexible and can convey much more information.

Overview

In this chapter, you'll learn how to:

- Inspect the data and its properties

- Compute descriptive statistics
- Create data visualizations inside and outside the command line

This chapter starts with the following files:

```
$ cd /data/ch07  
$ l  
total 104K  
-rw-r--r-- 1 dst dst 125 Jun 29 14:30 datatypes.csv  
-rw-r--r-- 1 dst dst 7.8K Jun 29 14:30 tips.csv  
-rw-r--r-- 1 dst dst 83K Jun 29 14:30 venture.csv  
-rw-r--r-- 1 dst dst 4.6K Jun 29 14:30 venture-wide.csv
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Inspecting Data and Its Properties

In this section I'll demonstrate how to inspect your dataset and its properties. Because the upcoming visualization and modeling techniques expect the data to be in a rectangular shape, I'll assume that the data is in CSV format. You can use the techniques described in [Chapter 5](#) to convert your data to CSV if necessary.

For simplicity's sake, I'll also assume that your data has a header. In the first subsection, I'll show you a way to determine whether that's the case. Once you know you have a header, you can continue answering the following questions:

- How many data points and features does the dataset have?
- What does the raw data look like?
- What kind of features does the dataset have?
- Can some of these features be treated as categorical?

Header or Not, Here I Come

You can check whether your file has a header by printing the first few lines using `head`:

```
$ head -n 5 venture.csv
FREQ,TIME_FORMAT,TIME_PERIOD,EXPEND,UNIT,GEO,OBS_STATUS,OBS_VALUE,FREQ_DESC,TIME
_FORMAT_DESC,TIME_PERIOD_DESC,OBS_STATUS_DESC,EXPEND_DESC,UNIT_DESC,GEO_DESC
A,P1Y,2015,INV_VEN,PC_GDP,CZ,,0.002,Annual,Annual,Year 2015,No data,"Venture
cap
ital investment (seed, start-up and later stage) ",Percentage of GDP,Czechia
A,P1Y,2007,INV_VEN,PC_GDP,DE,,0.034,Annual,Annual,Year 2007,No data,"Venture
cap
ital investment (seed, start-up and later stage) ",Percentage of GDP,Germany
A,P1Y,2008,INV_VEN,PC_GDP,DE,,0.039,Annual,Annual,Year 2008,No data,"Venture
cap
ital investment (seed, start-up and later stage) ",Percentage of GDP,Germany
A,P1Y,2009,INV_VEN,PC_GDP,DE,,0.029,Annual,Annual,Year 2009,No data,"Venture
cap
ital investment (seed, start-up and later stage) ",Percentage of GDP,Germany
```

If the lines wrap around, add line numbers using `nl`:

```
$ head -n 3 venture.csv | nl
1
FREQ,TIME_FORMAT,TIME_PERIOD,EXPEND,UNIT,GEO,OBS_STATUS,OBS_VALUE,FREQ_D
ESC,TIME_FORMAT_DESC,TIME_PERIOD_DESC,OBS_STATUS_DESC,EXPEND_DESC,UNIT_DESC,GE
O_
DESC
2 A,P1Y,2015,INV_VEN,PC_GDP,CZ,,0.002,Annual,Annual,Year 2015,No
data,"Ven
ture capital investment (seed, start-up and later stage) ",Percentage of
GDP,Cze
chia
3 A,P1Y,2007,INV_VEN,PC_GDP,DE,,0.034,Annual,Annual,Year 2007,No
data,"Ven
ture capital investment (seed, start-up and later stage) ",Percentage of
GDP,Ger
many
```

Alternatively, you can use `trim`:

```
$ < venture.csv trim 5
FREQ,TIME_FORMAT,TIME_PERIOD,EXPEND,UNIT,GEO,OBS_STATUS,OBS_VALUE,FREQ_DESC,TI
M...
A,P1Y,2015,INV_VEN,PC_GDP,CZ,,0.002,Annual,Annual,Year 2015,No data,"Venture
ca...
A,P1Y,2007,INV_VEN,PC_GDP,DE,,0.034,Annual,Annual,Year 2007,No data,"Venture
ca...
A,P1Y,2008,INV_VEN,PC_GDP,DE,,0.039,Annual,Annual,Year 2008,No data,"Venture
ca...
A,P1Y,2009,INV_VEN,PC_GDP,DE,,0.029,Annual,Annual,Year 2009,No data,"Venture
ca...
... with 536 more lines
```

In this case, it's clear that the first line is a header because it contains only uppercase names, and subsequent lines contain numbers. This is quite a subjective process, and it's up to you to decide whether the first line is a header or is already the first data point. When the dataset contains no header, you're best off using the `header` tool (discussed in [Chapter 5](#)) to correct that.

Inspect All the Data

If you want to inspect the raw data at your own pace, then it's probably not a good idea to use `cat`, because then all the data will be printed in one go. I recommend using `less`,¹ which allows you to interactively inspect your data in the command line. You can prevent long lines (as with `venture.csv`) from wrapping by specifying the `-S` option:

```
$ less -S venture.csv

FREQ,TIME_FORMAT,TIME_PERIOD,EXPEND,UNIT,GEO,OBS_STATUS,OBS_VALUE,FREQ_DESC,TI
M>
A,P1Y,2015,INV_VEN,PC_GDP,CZ,,0.002,Annual,Annual,Year 2015,No data,"Venture
ca>
A,P1Y,2007,INV_VEN,PC_GDP,DE,,0.034,Annual,Annual,Year 2007,No data,"Venture
ca>
A,P1Y,2008,INV_VEN,PC_GDP,DE,,0.039,Annual,Annual,Year 2008,No data,"Venture
ca>
A,P1Y,2009,INV_VEN,PC_GDP,DE,,0.029,Annual,Annual,Year 2009,No data,"Venture
ca>
A,P1Y,2010,INV_VEN,PC_GDP,DE,,0.029,Annual,Annual,Year 2010,No data,"Venture
```

```

ca>
A,P1Y,2011,INV_VEN,PC_GDP,DE,,0.029,Annual,Annual,Year 2011,No data,"Venture
ca>
A,P1Y,2012,INV_VEN,PC_GDP,DE,,0.021,Annual,Annual,Year 2012,No data,"Venture
ca>
A,P1Y,2013,INV_VEN,PC_GDP,DE,,0.023,Annual,Annual,Year 2013,No data,"Venture
ca>
A,P1Y,2014,INV_VEN,PC_GDP,DE,,0.021,Annual,Annual,Year 2014,No data,"Venture
ca>
A,P1Y,2015,INV_VEN,PC_GDP,DE,,0.025,Annual,Annual,Year 2015,No data,"Venture
ca>
A,P1Y,2007,INV_VEN,PC_GDP,DK,,0.092,Annual,Annual,Year 2007,No data,"Venture
ca>
A,P1Y,2008,INV_VEN,PC_GDP,DK,,0.074,Annual,Annual,Year 2008,No data,"Venture
ca>
A,P1Y,2009,INV_VEN,PC_GDP,DK,,0.051,Annual,Annual,Year 2009,No data,"Venture
ca>
A,P1Y,2010,INV_VEN,PC_GDP,DK,,0.059,Annual,Annual,Year 2010,No data,"Venture
ca>
:

```

The greater-than signs on the right indicate that you can scroll horizontally. You can scroll up and down by pressing the up and down arrow keys. Press the space bar to scroll down an entire screen. Scrolling horizontally is done by pressing the left and right arrow keys. Press g and G to go to the start and the end of the file, respectively. Quitting `less` is done by pressing q. The manual page lists all the available key bindings.

One advantage of `less` is that it does not load the entire file into memory, which means it's fast even for viewing large files.

Feature Names and Data Types

The column (or feature) names may indicate the meaning of the feature. You can use the following `head` and `tr` combo for this:

```
$ < venture.csv head -n 1 | tr , '\n'
FREQ
TIME_FORMAT
TIME_PERIOD
EXPEND
UNIT
GEO
```

```
OBS_STATUS  
OBS_VALUE  
FREQ_DESC  
TIME_FORMAT_DESC  
TIME_PERIOD_DESC  
OBS_STATUS_DESC  
EXPEND_DESC  
UNIT_DESC  
GEO_DESC
```

This basic command assumes that the file is delimited by commas. A more robust approach is to use `csvcut`:

```
$ csvcut -n venture.csv  
1: FREQ  
2: TIME_FORMAT  
3: TIME_PERIOD  
4: EXPEND  
5: UNIT  
6: GEO  
7: OBS_STATUS  
8: OBS_VALUE  
9: FREQ_DESC  
10: TIME_FORMAT_DESC  
11: TIME_PERIOD_DESC  
12: OBS_STATUS_DESC  
13: EXPEND_DESC  
14: UNIT_DESC  
15: GEO_DESC
```

You can go a step further than just printing the column names. Besides the names of the columns, it would be very useful to know what type of values each column contains, such as a string of characters, a numerical value, or a date. Assume that you have the following toy dataset:

```
$ bat -A datatype.csv  
-----  
--  
| File: datatype.csv  
-----  
--  
1 | a,b,c,d,e,f ↵  
2 | 1,0.0,TRUE,"""Yes!""",2011-11-11•11:00,2012-09-08 ↵  
3 | 42,3.1415,TRUE,"OK, • good",2014-09-15,12/6/70 ↵
```

```
4 | 66,,False,2198,,LF
```

Which `csvlook` interprets as follows:

```
$ csvlook datatypes.csv
```

a	b	c	d	e	f
1	0.000...	False	"Yes!"	2011-11-11 11:00:00	2012-09-08
42	3.142...	True	OK, good	2014-09-15 00:00:00	1970-12-06
66		False	2198		

I have already used `csvsql` in [Chapter 5](#) to execute SQL queries directly on CSV data. When no command-line arguments are passed, it generates the SQL statement that would be needed if you were to insert this data into an actual database. You can also use the output to inspect what the inferred column types are. If a column has the NOT NULL string printed after the data type, then that column contains no missing values:

```
$ csvsql datatypes.csv
```

```
CREATE TABLE datatypes (
    a DECIMAL NOT NULL,
    b DECIMAL,
    c BOOLEAN NOT NULL,
    d VARCHAR NOT NULL,
    e TIMESTAMP,
    f DATE
);
```

This output is especially useful when you use other tools within the `csvkit` suite, such as `csvgrep`, `csvsort`, and `csvsql`. For `venture.csv`, the columns are inferred as follows:

```
$ csvsql venture.csv
```

```
CREATE TABLE venture (
    "FREQ" VARCHAR NOT NULL,
    "TIME_FORMAT" VARCHAR NOT NULL,
    "TIME_PERIOD" DECIMAL NOT NULL,
    "EXPEND" VARCHAR NOT NULL,
    "UNIT" VARCHAR NOT NULL,
```

```

    "GEO" VARCHAR NOT NULL,
    "OBS_STATUS" BOOLEAN,
    "OBS_VALUE" DECIMAL NOT NULL,
    "FREQ_DESC" VARCHAR NOT NULL,
    "TIME_FORMAT_DESC" VARCHAR NOT NULL,
    "TIME_PERIOD_DESC" VARCHAR NOT NULL,
    "OBS_STATUS_DESC" VARCHAR NOT NULL,
    "EXPEND_DESC" VARCHAR NOT NULL,
    "UNIT_DESC" VARCHAR NOT NULL,
    "GEO_DESC" VARCHAR NOT NULL
);

```

Unique Identifiers, Continuous Variables, and Factors

Knowing the data type of each feature is not enough. It's also essential to know what each feature represents. Having knowledge about the domain is very useful here, but we may also get some context by looking at the data itself.

Both a string and an integer could be a unique identifier or could represent a category. In the latter case, this could be used to assign a color to your visualization. But if an integer denotes, say, a postal code, then it doesn't make sense to compute the average.

To determine whether a feature should be treated as a unique identifier or a categorical variable, you could count the number of unique values for a specific column:

```

$ wc -l tips.csv
245 tips.csv

$ < tips.csv csvcut -c day | header -d | sort | uniq | wc -l
4

```

You can use `csvstat`,² which is part of `csvkit`, to get the number of unique values for each column:

```

$ csvstat tips.csv --unique
1. bill: 229
2. tip: 123
3. sex: 2

```

```

4. smoker: 2
5. day: 4
6. time: 2
7. size: 6

$ csvstat venture.csv --unique
1. FREQ: 1
2. TIME_FORMAT: 1
3. TIME_PERIOD: 9
4. EXPEND: 1
5. UNIT: 3
6. GEO: 20
7. OBS_STATUS: 1
8. OBS_VALUE: 286
9. FREQ_DESC: 1
10. TIME_FORMAT_DESC: 1
11. TIME_PERIOD_DESC: 9
12. OBS_STATUS_DESC: 1
13. EXPEND_DESC: 1
14. UNIT_DESC: 3
15. GEO_DESC: 20

```

If there's only one unique value (such as with *OBS_STATUS*), then there's a chance that you can discard that column because it doesn't provide any value. If you wanted to automatically discard all such columns, then you could use the following pipeline:

```

$ < venture.csv csvcut -C $(①
>   csvstat venture.csv --unique | ②
>   grep ': 1$' | ③
>   cut -d. -f 1 | ④
>   tr -d ' ' | ⑤
>   paste -sd, ⑥
> ) | trim ⑦
TIME_PERIOD,UNIT,GEO,OBS_VALUE,TIME_PERIOD_DESC,UNIT_DESC,GEO_DESC
2015,PC_GDP,CZ,0.002,Year 2015,Percentage of GDP,Czechia
2007,PC_GDP,DE,0.034,Year 2007,Percentage of GDP,Germany
2008,PC_GDP,DE,0.039,Year 2008,Percentage of GDP,Germany
2009,PC_GDP,DE,0.029,Year 2009,Percentage of GDP,Germany
2010,PC_GDP,DE,0.029,Year 2010,Percentage of GDP,Germany
2011,PC_GDP,DE,0.029,Year 2011,Percentage of GDP,Germany
2012,PC_GDP,DE,0.021,Year 2012,Percentage of GDP,Germany
2013,PC_GDP,DE,0.023,Year 2013,Percentage of GDP,Germany
2014,PC_GDP,DE,0.021,Year 2014,Percentage of GDP,Germany
... with 531 more lines

```

- ❶ The `-C` option deselects columns given their locations (or names), which is provided with command substitution.
- ❷ Obtain the number of unique values for each column in `venture.csv`.
- ❸ Keep only the columns that contain one unique value.
- ❹ Extract the column location.
- ❺ Trim any white space.
- ❻ Put all column locations on one comma-separated line.
- ❼ Show only the first 10 lines.

Having said that, I'm going to keep those columns for now.

Generally speaking, if the number of unique values is low compared to the total number of rows, then that feature might be treated as a categorical one (such as `GEO` in the case of `venture.csv`). If the number is equal to the number of rows, it might be a unique identifier, but it might also be a numerical value. There's only one way to find out: we need to go deeper.

Computing Descriptive Statistics

In this section, we're going to use `csvstat` and `rush` to compute various descriptive statistics.

Column Statistics

The command-line tool `csvstat` gives a lot of information. For each feature (column), it shows:

- The data type
- Whether it has any missing values (nulls)
- The number of unique values
- Various descriptive statistics (minimum, maximum, sum, mean, standard deviation, and median) for those features for which it is appropriate

Invoke `csvstat` as follows:

```
$ csvstat venture.csv | trim 32
1. "FREQ"

  Type of data:      Text
  Contains null values: False
  Unique values:      1
  Longest value:      1 characters
  Most common values: A (540x)

2. "TIME_FORMAT"

  Type of data:      Text
  Contains null values: False
  Unique values:      1
  Longest value:      3 characters
  Most common values: P1Y (540x)

3. "TIME_PERIOD"

  Type of data:      Number
  Contains null values: False
  Unique values:      9
  Smallest value:     2,007
  Largest value:      2,015
  Sum:                1,085,940
  Mean:               2,011
  Median:              2,011
  StDev:                2.584
  Most common values: 2,015 (60x)
                      2,007 (60x)
                      2,008 (60x)
                      2,009 (60x)
```

```
2,010 (60x)
... with 122 more lines
```

I'm showing only the first 32 lines because this produces a lot of output. You might want to pipe this through `less`. If you're only interested in a specific statistic, you can also use one of the following options:

- `--max` (maximum)
- `--min` (minimum)
- `--sum` (sum)
- `--mean` (mean)
- `--median` (median)
- `--stdev` (standard deviation)
- `--nulls` (whether a column contains nulls)
- `--unique` (unique values)
- `--freq` (frequent values)
- `--len` (maximum value length)

For example:

```
$ csvstat venture.csv --freq | trim
1. FREQ: { "A": 540 }
2. TIME_FORMAT: { "P1Y": 540 }
3. TIME_PERIOD: { "2015": 60, "2007": 60, "2008": 60, "2009": 60, "2010": 60
}
4. EXPEND: { "INV_VEN": 540 }
5. UNIT: { "PC_GDP": 180, "NR_COMP": 180, "MIO_EUR": 180 }
6. GEO: { "CZ": 27, "DE": 27, "DK": 27, "EL": 27, "ES": 27 }
7. OBS_STATUS: { "None": 540 }
8. OBS_VALUE: { "0": 28, "1": 19, "2": 14, "0.002": 10, "0.034": 7 }
9. FREQ_DESC: { "Annual": 540 }
10. TIME_FORMAT_DESC: { "Annual": 540 }
... with 5 more lines
```

You can select a subset of features with the `-c` option, which accepts both integers and column names:

```
$ csvstat venture.csv -c 3,GEO  
3. "TIME_PERIOD"
```

Type of data:	Number
Contains null values:	False
Unique values:	9
Smallest value:	2,007
Largest value:	2,015
Sum:	1,085,940
Mean:	2,011
Median:	2,011
StDev:	2.584
Most common values:	2,015 (60x) 2,007 (60x) 2,008 (60x) 2,009 (60x) 2,010 (60x)

6. "GEO"

Type of data:	Text
Contains null values:	False
Unique values:	20
Longest value:	2 characters
Most common values:	CZ (27x) DE (27x) DK (27x) EL (27x) ES (27x)

Row count: 540

TIP

Keep in mind that `csvstat`, just like `csvsql`, employs heuristics to determine the data type and therefore may not always get it right. I encourage you to always do a manual inspection, as discussed in the previous subsection. Moreover, even though the type may be a string or an integer, that doesn't say anything about how it should be used.

As a nice extra, `csvstat` outputs, at the very end, the number of data points (rows). Newlines and commas inside values are handled correctly. To see only that last line, you can use `tail`. Alternatively, you can use `xsv`, which returns only the actual number of rows:

```
$ csvstat venture.csv | tail -n 1  
Row count: 540
```

```
$ xsv count venture.csv  
540
```

Note that using either of these two options is different from using `wc -l`, which counts the number of newlines (and therefore also counts the header).

R One-Liners on the Shell

In this section I'd like to discuss the command-line tool `rush`, which enables you to leverage the statistical programming environment R³ directly from the command line. Before I explain what `rush` does and why it exists, let's talk a bit about R itself.

R is a very powerful statistical software package for doing data science. It's an interpreted programming language, has an extensive collection of packages, and offers its own REPL, which, similar to the command line, allows you to play with your data. Note that once you start R, you're in an interactive session that is separated from the Unix command line.

Imagine that you have a CSV file called `tips.csv`, and you would like compute the tip percentage and save the result. To accomplish this in R, you would first run R:

```
$ R --quiet ❶  
>
```

❶ I use the `--quiet` option here to suppress the rather long startup message.

And then run the following code:

```
> library(tidyverse)                                     ❶
— Attaching packages ——————— tidyverse 1.3.0
—
✓ ggplot2 3.3.3     ✓ purrr   0.3.4
✓ tibble  3.0.6     ✓ dplyr    1.0.4
✓ tidyr   1.1.2     ✓ stringr 1.4.0
✓ readr   1.4.0     ✓ forcats 0.5.1
— Conflicts ——————— tidyverse_conflicts()
—
✖ dplyr::filter() masks stats::filter()
✖ dplyr::lag()   masks stats::lag()
> df <- read_csv("tips.csv")                         ❷

— Column specification ———————
cols(
  bill = col_double(),
  tip = col_double(),
  sex = col_character(),
  smoker = col_character(),
  day = col_character(),
  time = col_character(),
  size = col_double()
)

> df <- mutate(df, percent = tip / bill * 100)      ❸
> write_csv(df, "percent.csv")                        ❹
> q("no")                                            ❺

$
```

- ❶ Load any required packages.
- ❷ Read in the CSV file and assign it to a variable.
- ❸ Compute the new column *percent*.
- ❹ Save the result to disk.

- ⑤ Exit R.

Afterward, you can continue with the saved file *percent.csv* on the command line:

```
$ < percent.csv trim 5  
bill,tip,sex,smoker,day,time,size,percent  
16.99,1.01,Female,No,Sun,Dinner,2,5.9446733372572105  
10.34,1.66,Male,No,Sun,Dinner,3,16.054158607350097  
21.01,3.5,Male,No,Sun,Dinner,3,16.658733936220845  
23.68,3.31,Male,No,Sun,Dinner,2,13.97804054054054  
... with 240 more lines
```

Note that only the third line is associated with what you specifically want to accomplish. The other lines are necessary boilerplate. Typing in this boilerplate in order to accomplish something simple is cumbersome and breaks your workflow. Sometimes, you want to do only one or two things at a time to your data. Wouldn't it be great if you could harness the power of R and use it from the command line?

This is where `rush` comes in. Let's perform the same task as before, but now using `rush`:

```
$ rm percent.csv  
  
$ rush run -t 'mutate(df, percent = tip / bill * 100)' tips.csv > percent.csv  
  
$ < percent.csv trim 5  
bill,tip,sex,smoker,day,time,size,percent  
16.99,1.01,Female,No,Sun,Dinner,2,5.9446733372572105  
10.34,1.66,Male,No,Sun,Dinner,3,16.054158607350097  
21.01,3.5,Male,No,Sun,Dinner,3,16.658733936220845  
23.68,3.31,Male,No,Sun,Dinner,2,13.97804054054054  
... with 240 more lines
```

These small one-liners are possible because `rush` takes care of all the boilerplate. In this case, I'm using the `run` subcommand. There's also the `plot` subcommand, which I'll use in the next section to produce data visualizations quickly. If you're passing in any input data, then by default,

`rush` assumes that it's in CSV format with a header and a comma as the delimiter. Moreover, the column names are sanitized so that they are easier to work with. You can override these defaults using the `--no-header` (or `-H`), `--delimiter` (or `-d`), and `--no-clean-names` (or `-C`) options, respectively. The help gives a good overview of the available options for the `run` subcommand:

```
$ rush run --help
rush: Run an R expression

Usage:
  rush run [options] <expression> [--] [<file>...]

Reading options:
  -d, --delimiter <str>      Delimiter [default: ,].
  -C, --no-clean-names        No clean names.
  -H, --no-header             No header.

Setup options:
  -l, --library <name>       Libraries to load.
  -t, --tidyverse              Enter the Tidyverse.

Saving options:
  --dpi <str|int>            Plot resolution [default: 300].
  --height <int>              Plot height.
  -o, --output <str>          Output file.
  --units <str>                Plot size units [default: in].
  -w, --width <int>            Plot width.

General options:
  -n, --dry-run               Only print generated script.
  -h, --help                  Show this help.
  -q, --quiet                 Be quiet.
  --seed <int>                Seed random number generator.
  -v, --verbose                Be verbose.
  --version                  Show version.
```

Under the hood, `rush` generates an R script and subsequently executes it. You can view this generated script by specifying the `--dry-run` (or `-n`) option:

```
$ rush run -n --tidyverse 'mutate(df, percent = tip / bill * 100)' tips.csv
#!/usr/bin/env Rscript
```

```

library(tidyverse)
library(glue)
df <- janitor::clean_names(readr::read_delim("tips.csv", delim = ",",
col_names
= TRUE))
mutate(df, percent = tip/bill * 100)

```

This generated script:

- Writes out the shebang (`#!/`; see [Chapter 4](#)) needed for running an R script from the command line
- Imports the tidyverse and glue packages
- Loads *tips.csv* as a data frame, cleans the column names, and assigns it to a variable `df`
- Runs the specified expression
- Prints the result to standard output

You could redirect this generated script to a file and easily turn it into a new command-line tool because of the shebang.

The output of `rush` doesn't have to be in CSV format per se. Here, I compute the mean tip percent, the maximum party size, the unique values of the *time* column, and the correlation between the *bill* and the *tip*. Finally, I extract an entire column (but show only the first 10 values):

```

$ < percent.csv rush run 'mean(df$percent)' -
16.0802581722505

$ < percent.csv rush run 'max(df$size)' -
6

$ < percent.csv rush run 'unique(df$time)' -
Dinner
Lunch

$ < percent.csv rush run 'cor(df$bill, df$tip)' -
0.675734109211365

$ < percent.csv rush run 'df$tip' - | trim

```

```
1.01
1.66
3.5
3.31
3.61
4.71
2
3.12
1.96
3.23
... with 234 more lines
```

That last dash means that `rush` should read from standard input.

So now if you want to do one or two things to your dataset with R, you can specify it as a one-liner and keep on working on the command line. All the knowledge that you already have about R can now be used from the command line. With `rush`, you can even create sophisticated visualizations, as I'll show you in the next section.

Creating Visualizations

In this section, I'm going to show you how to create data visualizations at the command line. Using `rush plot`, I'll be creating bar charts, scatter plots, and box plots. Before we dive in, though, I'd first like to explain how you can display your visualizations.

Displaying Images from the Command Line

Let's take the image `tips.png` as an example. [Figure 7-1](#) is a data visualization that was created using `rush` and the `tips.csv` dataset. (I'll explain the `rush` syntax in a moment.) I used the `display` tool to insert the image in the book, but if you run `display` you'll find that it doesn't work. That's because displaying images from the command line is actually quite tricky.

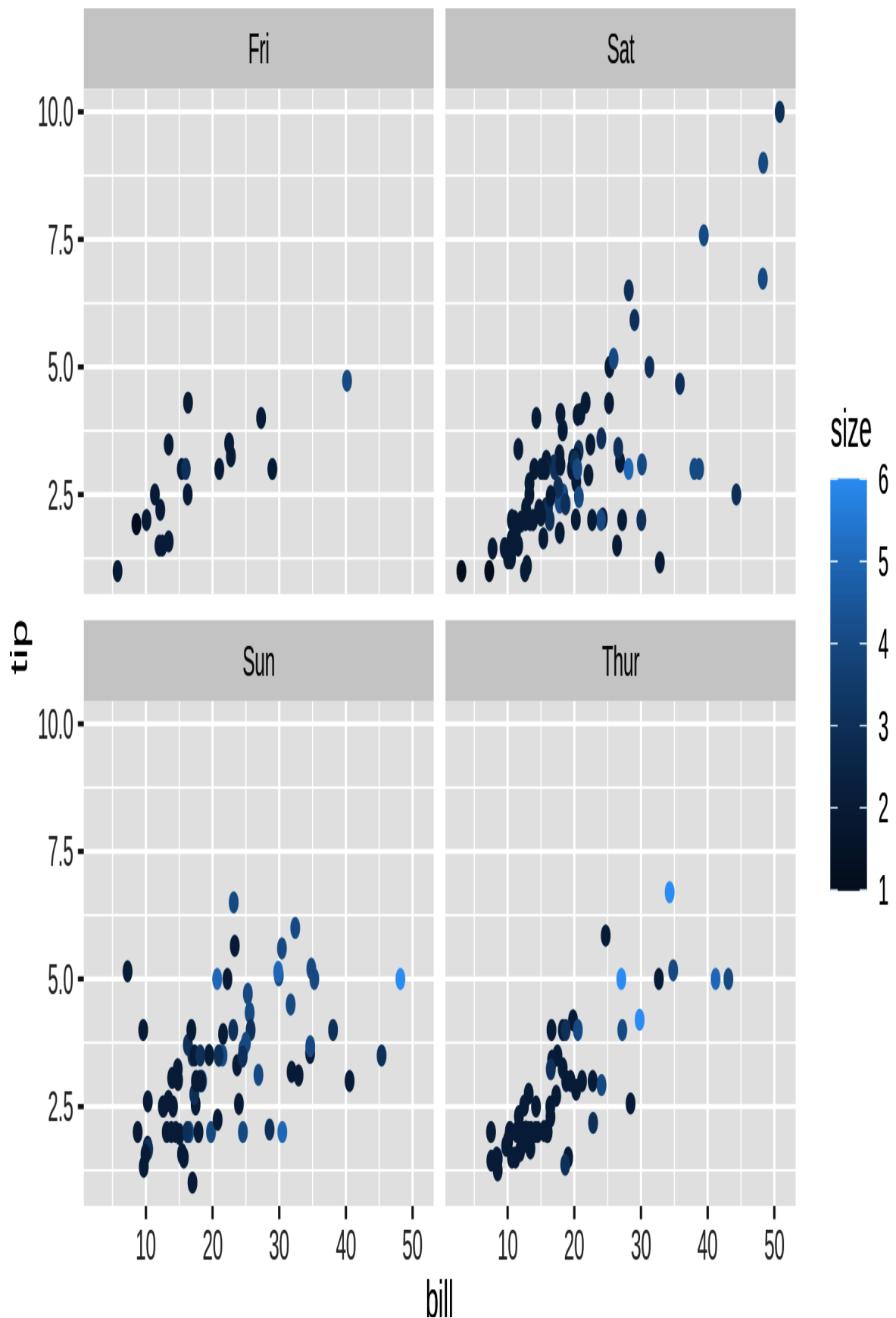


Figure 7-1. Displaying this image yourself can be tricky

Depending on your setup, there are different options available for displaying images. I know of four options, each with its own advantages and disadvantages: (1) as a textual representation, (2) as an inline image, (3) using an image viewer, and (4) using a browser. Let's go through them quickly.

Option 1 is to display the image inside the terminal, as shown at the top of [Figure 7-2](#).

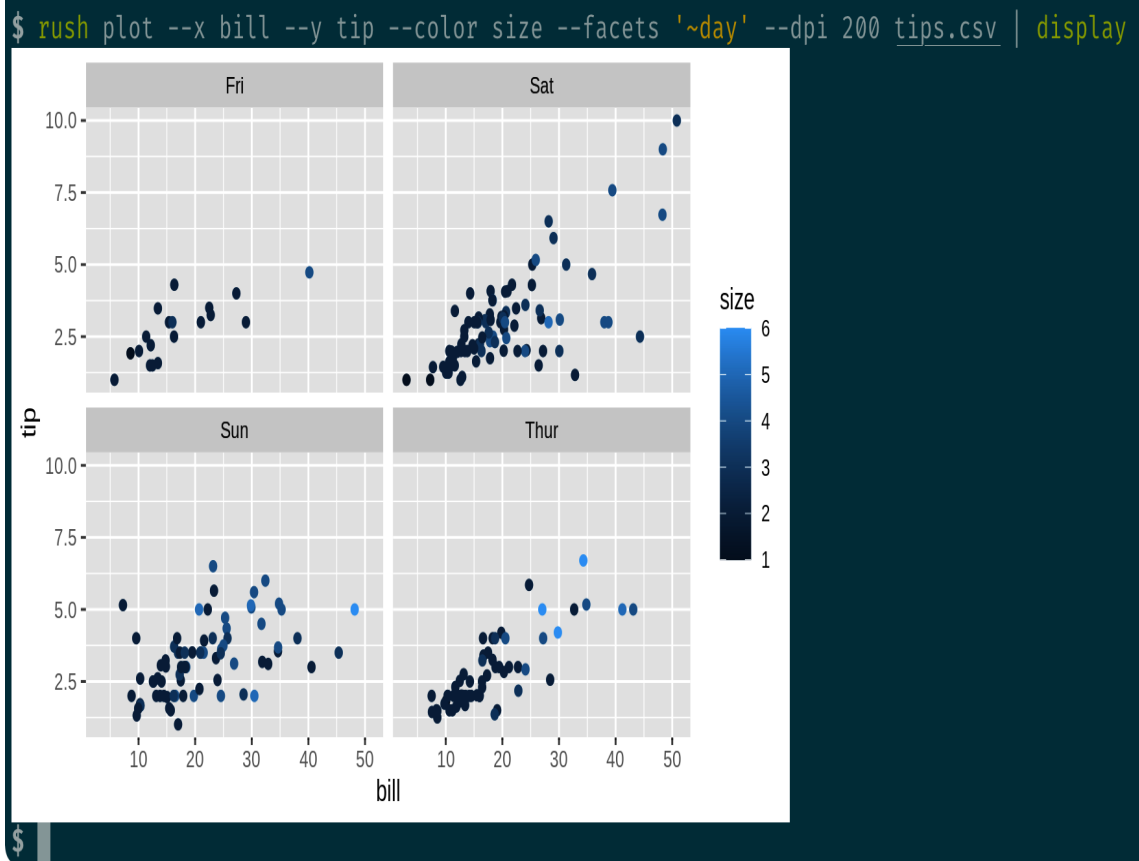
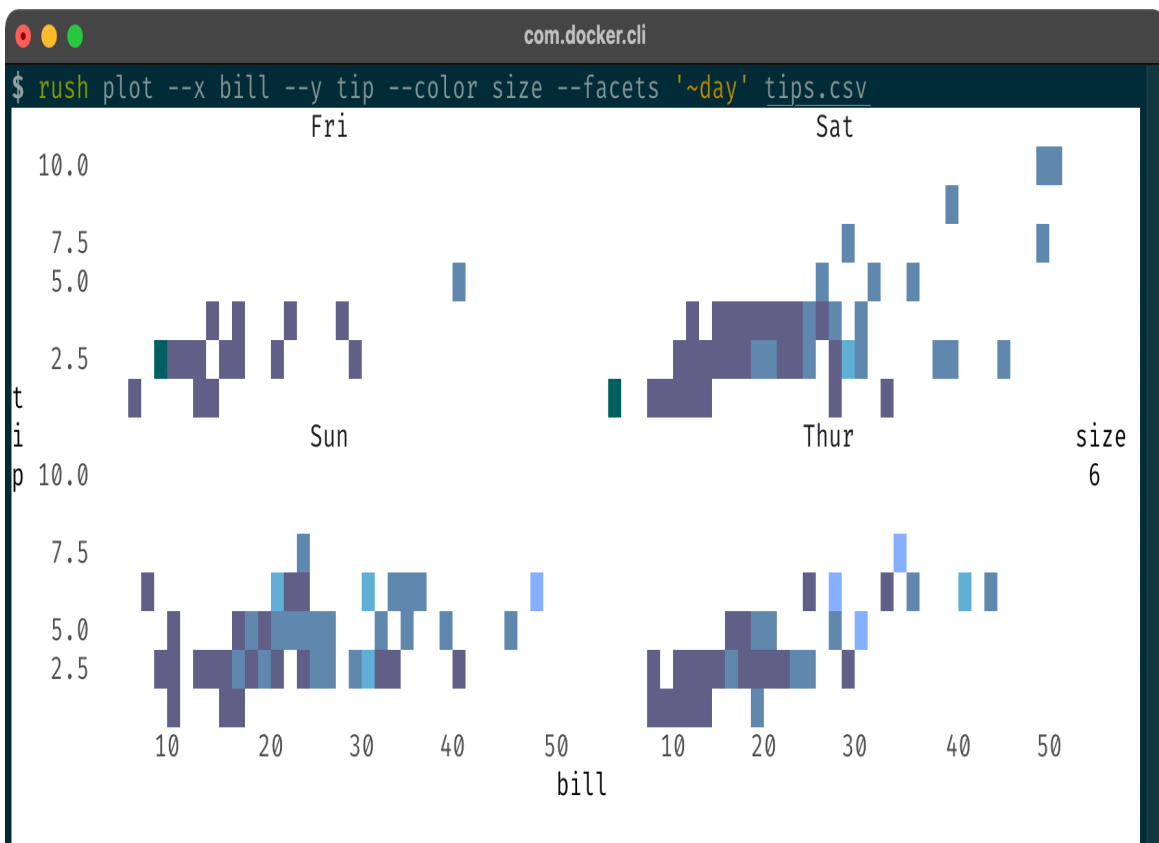
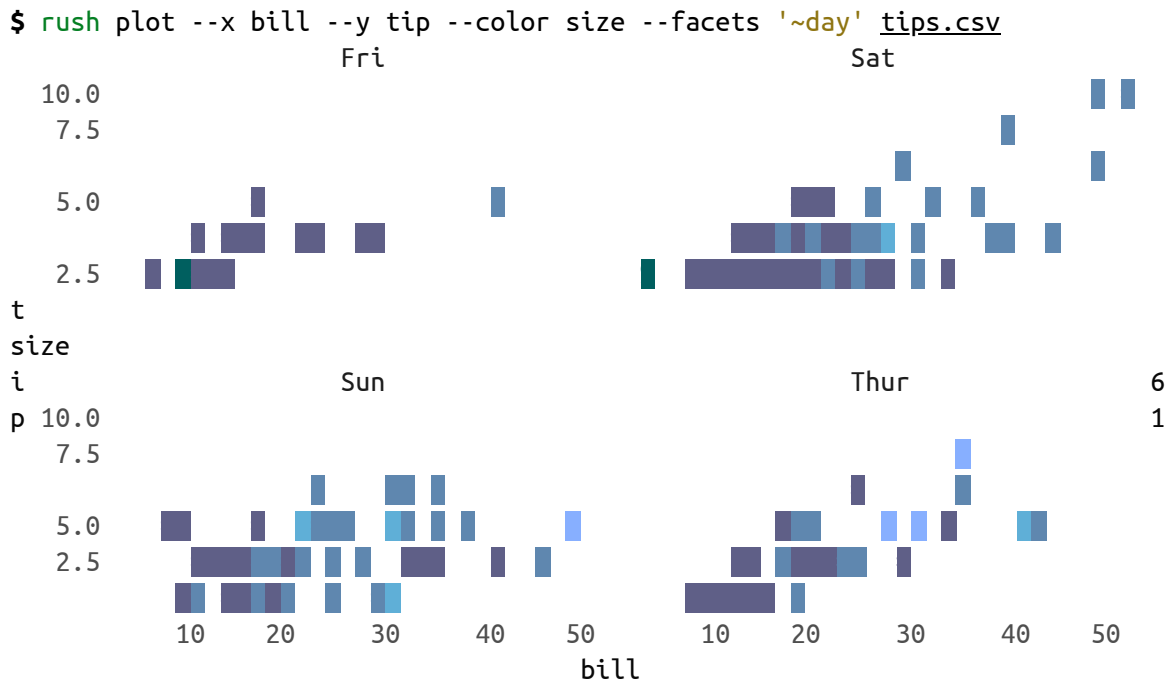


Figure 7-2. Displaying an image in the terminal via ASCII characters and ANSI escape sequences (top) and via the iTerm2 Inline Images Protocol (bottom)

This output is generated by `rush` when the standard output is not redirected to a file. It's based on ASCII characters and ANSI escape sequences, so it's available in every terminal. Depending on how you're reading this book, the output you get when you run the following code may or may not match the screenshot in [Figure 7-2](#):



If you see only ASCII characters, that means the medium on which you're reading this book doesn't support the ANSI escape sequences responsible for the colors. Fortunately, if you run the preceding command yourself, it will look just like the screenshot.

Option 2, as seen at the bottom of [Figure 7-2](#), also displays images inside the terminal. This is the iTerm2 terminal, which is only available for macOS and uses the [Inline Images Protocol](#) through a small script (which I have named `display`). This script is not included with the Docker image, but you can easily install it:

```
$ curl -s "https://iterm2.com/utilities/imgcat" > display && chmod u+x display
```

If you're not using iTerm2 on macOS, there might be other options available to display images inline. Please consult your favorite search engine.

Option 3 is to manually open the image (*tips.csv* in this example) in an image viewer. **Figure 7-3** shows, on the left, the file explorer (Finder) and image viewer (Preview) on macOS.

File: ch07

Name	Size	Kind
datatypes.csv	125 bytes	Comma-separated Value
tips.csv	8 KB	Comma-separated Value
tips.png	121 KB	PNG image
venture-wide.csv	5 KB	Comma-separated Value
venture.csv	84 KB	Comma-separated Value

```
$ rush plot --x bill --y tip --color size --facets '~day' tips.csv > tips.png
$ servewd
$ Serving HTTP on 0.0.0.0 port 8000 (http://0.0.0.0:8000/) ...
$ l
total 224K
-rw-r--r-- 1 dst dst 125 Apr  7 10:42 datatypes.csv
-rw-r--r-- 1 dst dst 7.8K Apr  7 10:42 tips.csv
-rw-r--r-- 1 dst dst 118K Jun  3 15:57 tips.png
-rw-r--r-- 1 dst dst 83K Apr  7 10:42 venture.csv
-rw-r--r-- 1 dst dst 4.6K Apr  7 10:42 venture-wide.csv
$
```

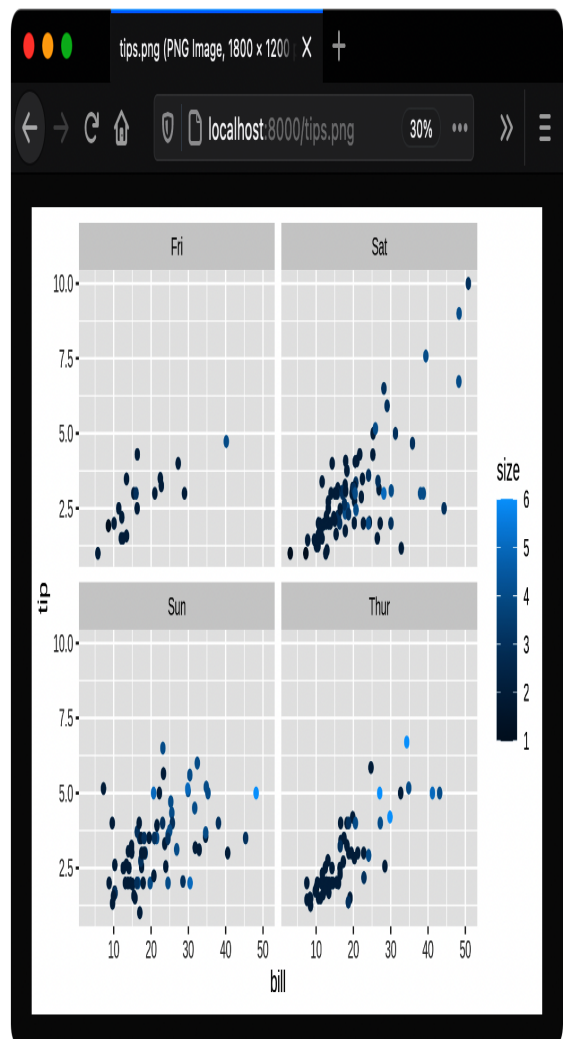
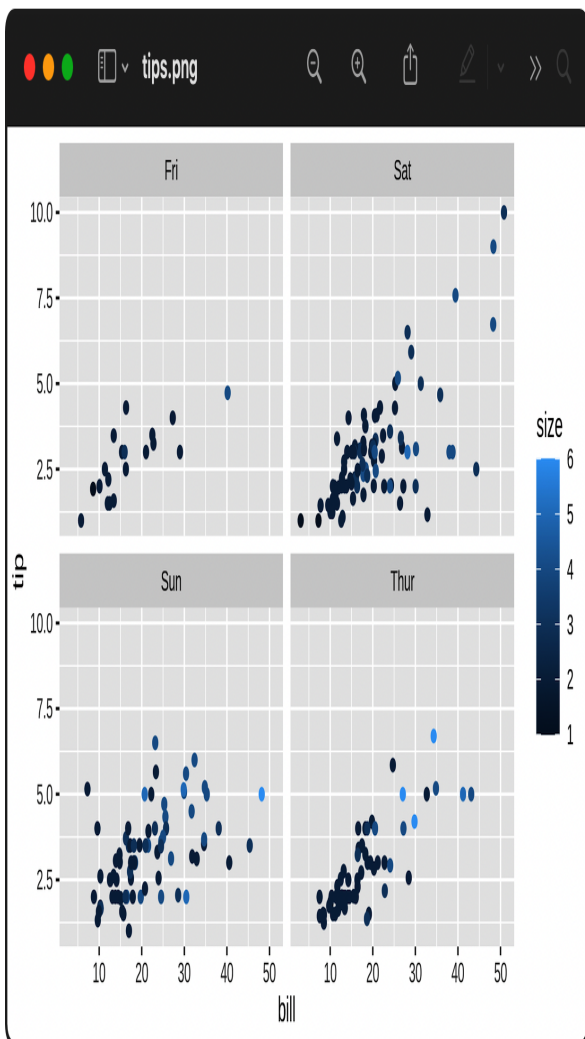


Figure 7-3. Displaying an image externally via a file explorer and an image viewer (left) and via a web server and a browser (right)

When you’re working locally, this option always works. When you’re working inside a Docker container, you can only access the generated image from your OS when you’ve mapped a local directory using the `-v` option. See [Chapter 2](#) for instructions on how to do this. An advantage of this option is that most image viewers automatically update the display when the image has changed, which allows for quick iterations as you fine-tune your visualization.

Option 4 is to open the image in a browser. The right side of [Figure 7-3](#) is a screenshot of Firefox showing <http://localhost:8000/tips.png>. You can use any browser for this, but you need two other prerequisites for it to work. First, you need to have made a port (port 8000 in this example) accessible on the Docker container using the `-p` option. (Again, see [Chapter 2](#) for instructions on how to do this.) Second, you need to start a web server. For this, the Docker container has a small tool called `servewd`,⁴ which serves the current working directory using Python:

```
$ bat $(which servewd)
-----
| File: /usr/bin/dsutils/servewd
-----
1 | #!/usr/bin/env bash
2 | ARGS="$@"
3 | python3 -m http.server ${ARGS} 2>/dev/null &
-----
```

You only need to run `servewd` once from a directory (`/data/`, for example), and it will happily run in the background. Once you’ve plotted something, you can visit `localhost:8000` in your browser and access the contents of that directory and all of its subdirectories. The default port is 8000, but you can change this by specifying it as an argument to `servewd`:

```
$ servewd 9999 > display
```

Just make sure that this port is accessible. Because `servewd` runs in the background, you need to stop it as follows:

```
$ pkill -f http.server
```

Option 4 can also work on a remote machine.

Now that we've covered four options for displaying images, let's move on to actually creating some.

Plotting in a Rush

When it comes to creating data visualizations, there's a plethora of options. Personally, I'm a staunch proponent of `ggplot2`, which is a visualization package for R. The underlying grammar of graphics is accompanied by a consistent API that allows you to quickly and iteratively create different types of beautiful data visualizations while rarely having to consult the documentation—a welcoming set of properties when exploring data.

We're not really in a rush, but we also don't want to fiddle too much with any single visualization. Moreover, we'd like to stay at the command line as much as possible. Luckily, we still have `rush`, which allows us to use `ggplot2` from the command line. The data visualization in [Figure 7-1](#) could have been created as follows:

```
$ rush run --library ggplot2 'ggplot(df, aes(x = bill, y = tip, color = size))  
+  
  geom_point() + facet_wrap(~day)' tips.csv > tips.png
```

However, as you may have noticed, I have used a very different command to create `tips.png`:

```
$ rush plot --x bill --y tip --color size --facets '~day' tips.csv > tips.png
```

While the syntax of `ggplot2` is relatively concise, especially considering the flexibility it offers, there's a shortcut to create basic plots quickly. This shortcut is available through the `plot` subcommand of `rush`, and it allows

you to create beautiful basic plots without needing to learn R and the grammar of graphics.

Under the hood, `rush plot` uses the function `qplot` from the `ggplot2` package. Here's the first part of `qplot`'s documentation:

```
$ R -q -e '?ggplot2::qplot' | trim 14
> ?ggplot2::qplot
qplot                               package:ggplot2                         R Documentation

Quick plot

Description:

  'qplot()' is a shortcut designed to be familiar if you're used to
  base 'plot()'. It's a convenient wrapper for creating a number of
  different types of plots using a consistent calling scheme. It's
  great for allowing you to produce plots quickly, but I highly
  recommend learning 'ggplot()' as it makes it easier to create
  complex graphics.

... with 108 more lines
```

I agree with this advice; once you're done reading this book, you'll find it worthwhile to learn `ggplot2`, especially if you want to upgrade any exploratory data visualizations into ones that are suitable for communication. For now, while we're at the command line, let's take that shortcut.

As [Figure 7-2](#) already showed, `rush plot` can create both graphical visualizations (consisting of pixels) and textual visualizations (consisting of ASCII characters and ANSI escape sequences) with the same syntax. When `rush` detects that its output has been piped to another command such as `display`, or redirected to a file such as `tips.png`, it will produce a graphical visualization; otherwise, it will produce a textual visualization.

Let's take a moment to read through the plotting and saving options of `rush plot`:

```
$ rush plot --help
rush: Quick plot
```

Usage:

`rush plot [options] [--] [<file>|-]`

Reading options:

<code>-d, --delimiter <str></code>	Delimiter [default: ,].
<code>-C, --no-clean-names</code>	No clean names.
<code>-H, --no-header</code>	No header.

Setup options:

<code>-l, --library <name></code>	Libraries to load.
<code>-t, --tidyverse</code>	Enter the Tidyverse.

Plotting options:

<code>--aes <key=value></code>	Additional aesthetics.
<code>-a, --alpha <name></code>	Alpha column.
<code>-c, --color <name></code>	Color column.
<code>--facets <formula></code>	Facet specification.
<code>-f, --fill <name></code>	Fill column.
<code>-g, --geom <geom></code>	Geometry [default: auto].
<code>--group <name></code>	Group column.
<code>--log <x y xy></code>	Variables to log transform.
<code>--margins</code>	Display marginal facets.
<code>--post <code></code>	Code to run after plotting.
<code>--pre <code></code>	Code to run before plotting.
<code>--shape <name></code>	Shape column.
<code>--size <name></code>	Size column.
<code>--title <str></code>	Plot title.
<code>-x, --x <name></code>	X column.
<code>--xlab <str></code>	X axis label.
<code>-y, --y <name></code>	Y column.
<code>--ylab <str></code>	Y axis label.
<code>-z, --z <name></code>	Z column.

Saving options:

<code>--dpi <str int></code>	Plot resolution [default: 300].
<code>--height <int></code>	Plot height.
<code>-o, --output <str></code>	Output file.
<code>--units <str></code>	Plot size units [default: in].
<code>-w, --width <int></code>	Plot width.

General options:

<code>-n, --dry-run</code>	Only print generated script.
<code>-h, --help</code>	Show this help.
<code>-q, --quiet</code>	Be quiet.
<code>--seed <int></code>	Seed random number generator.
<code>-v, --verbose</code>	Be verbose.
<code>--version</code>	Show version.

The most important options are the plotting options that take a `<name>` as an argument. For example, the `--x` option allows you to specify which column should be used to determine where *things* should be placed along the x-axis. The same holds for the `--y` option. The `--color` and `--fill` options are used to specify which column you want to use for coloring. You can probably guess what the `--size` and `--alpha` options are about. Other common options are explained throughout the sections as I create various visualizations. Note that for each visualization, I first show its textual representation (ASCII and ANSI characters) and then its visual representation (pixels).

Creating Bar Charts

Bar charts are especially useful for displaying the value counts of a categorical feature. Here's a textual visualization of the *time* feature in the *tips.csv* dataset:

```
$ rush plot --x time tips.csv
```

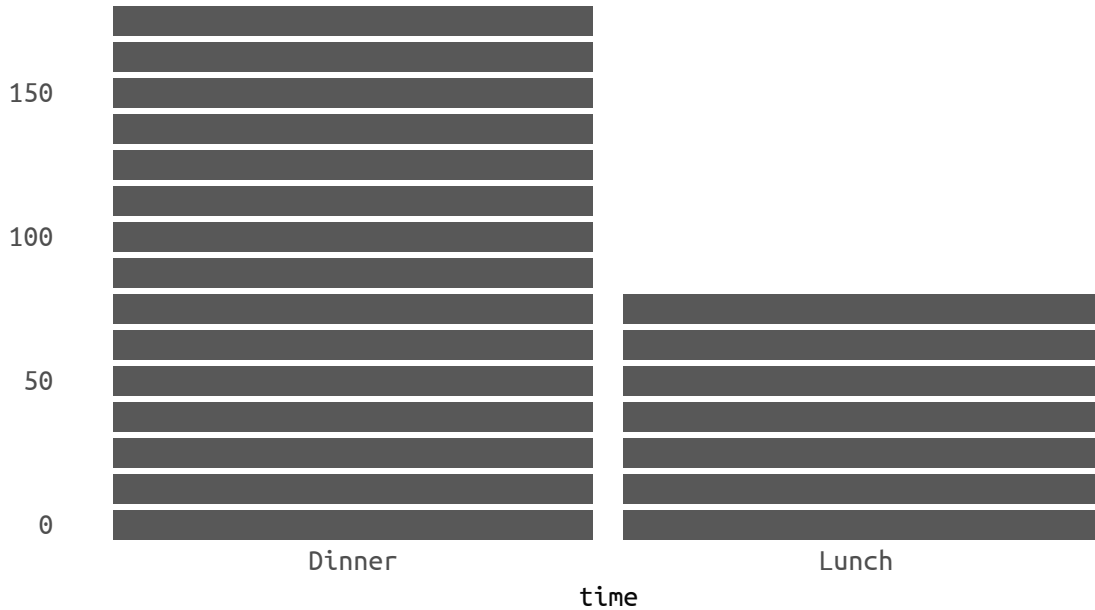


Figure 7-4 shows the graphical visualization, which is created by `rush plot` when the output is redirected to a file:

```
$ rush plot --x time tips.csv > plot-bar.png
```

```
$ display plot-bar.png
```

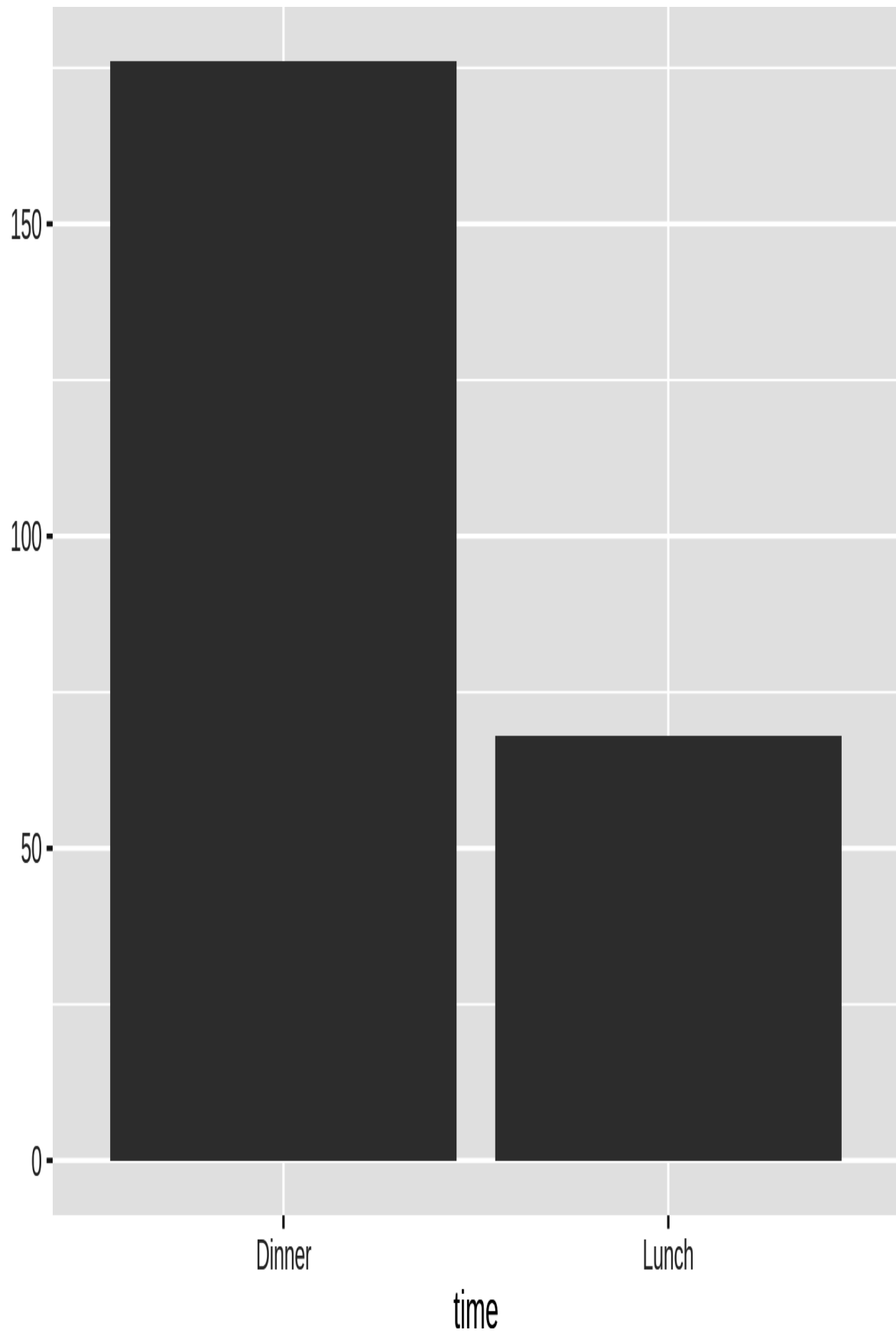
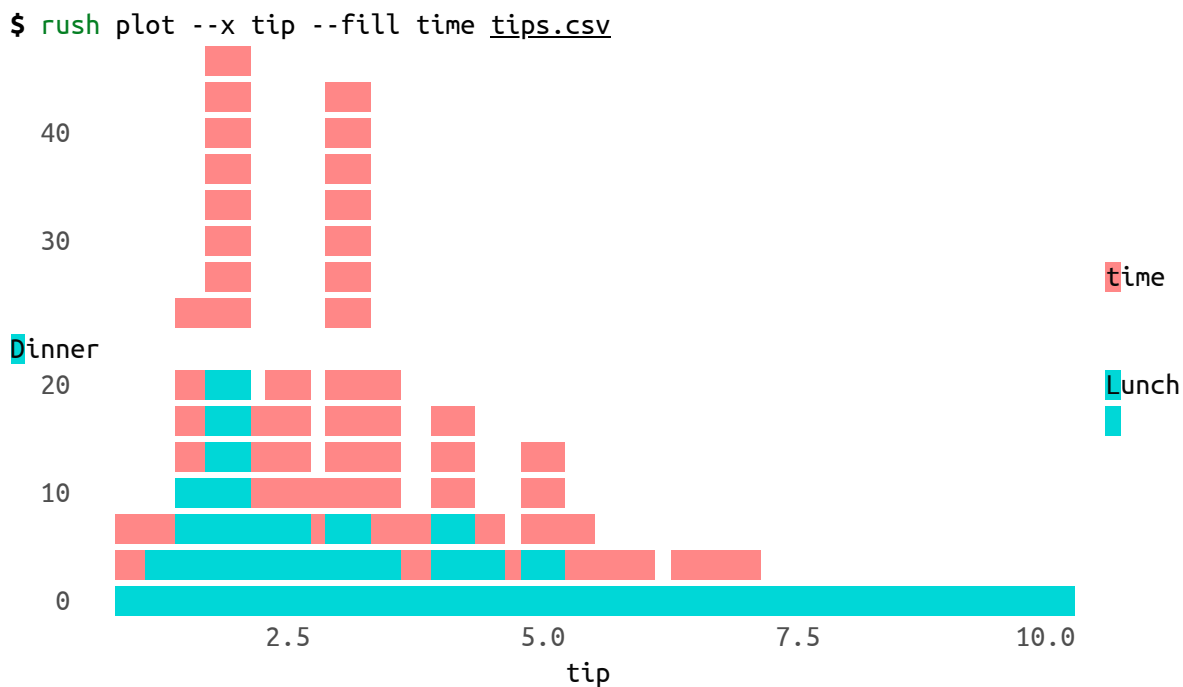


Figure 7-4. A bar chart

The conclusion we can draw from this bar chart is straightforward: there are more than twice as many data points for dinner than lunch.

Creating Histograms

The counts of a continuous variable can be visualized with a histogram. Here, I have used the *time* feature to set the fill color. As a result, `rush plot` conveniently creates a stacked histogram:



TIP

Allow me to demonstrate two syntax shortcuts that you may find useful. The two exclamation marks (!!) get replaced with the previous command. The exclamation mark and dollar sign (!\$) get replaced by the last part of the previous command, which is the filename `plot-histogram.png`. As you can see, the updated commands are first printed by the Z shell so you know exactly what it executes. These two shortcuts can save a lot of typing, but they're not easy to remember.

Figure 7-5 shows the graphical visualization:

```
$ !! > plot-histogram.png
rush plot --x tip --fill time tips.csv > plot-histogram.png

$ display !$
display plot-histogram.png
```

This histogram reveals that most tips are around \$2.50. Because the dinner and lunch groups are stacked on top of each other and show absolute counts, it's difficult to compare them. Perhaps a density plot can help with this.

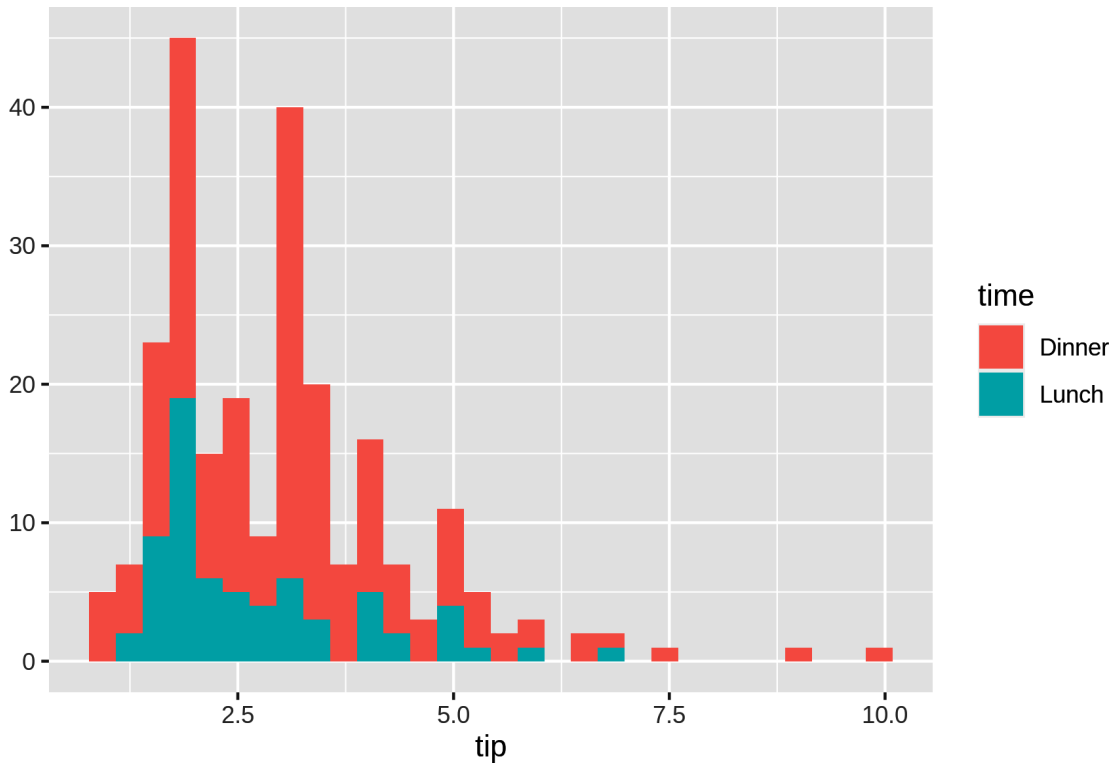
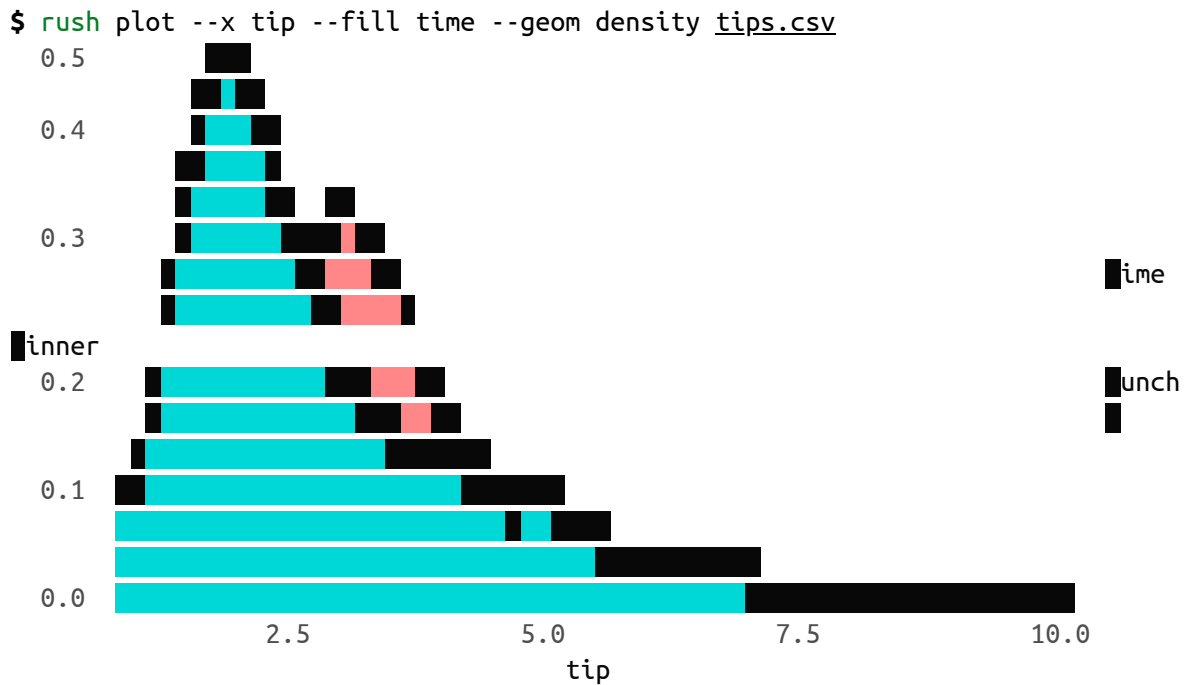


Figure 7-5. A histogram

Creating Density Plots

A density plot is useful for visualizing the distribution of a continuous variable. `rush plot` uses heuristics to determine the appropriate geometry, but you can override this with the `--geom` option:



In this case, the textual representation really shows its limitations when compared to the visual representation in [Figure 7-6](#):

```
$ rush plot --x tip --fill time --geom density tips.csv > plot-density.png
$ display plot-density.png
```

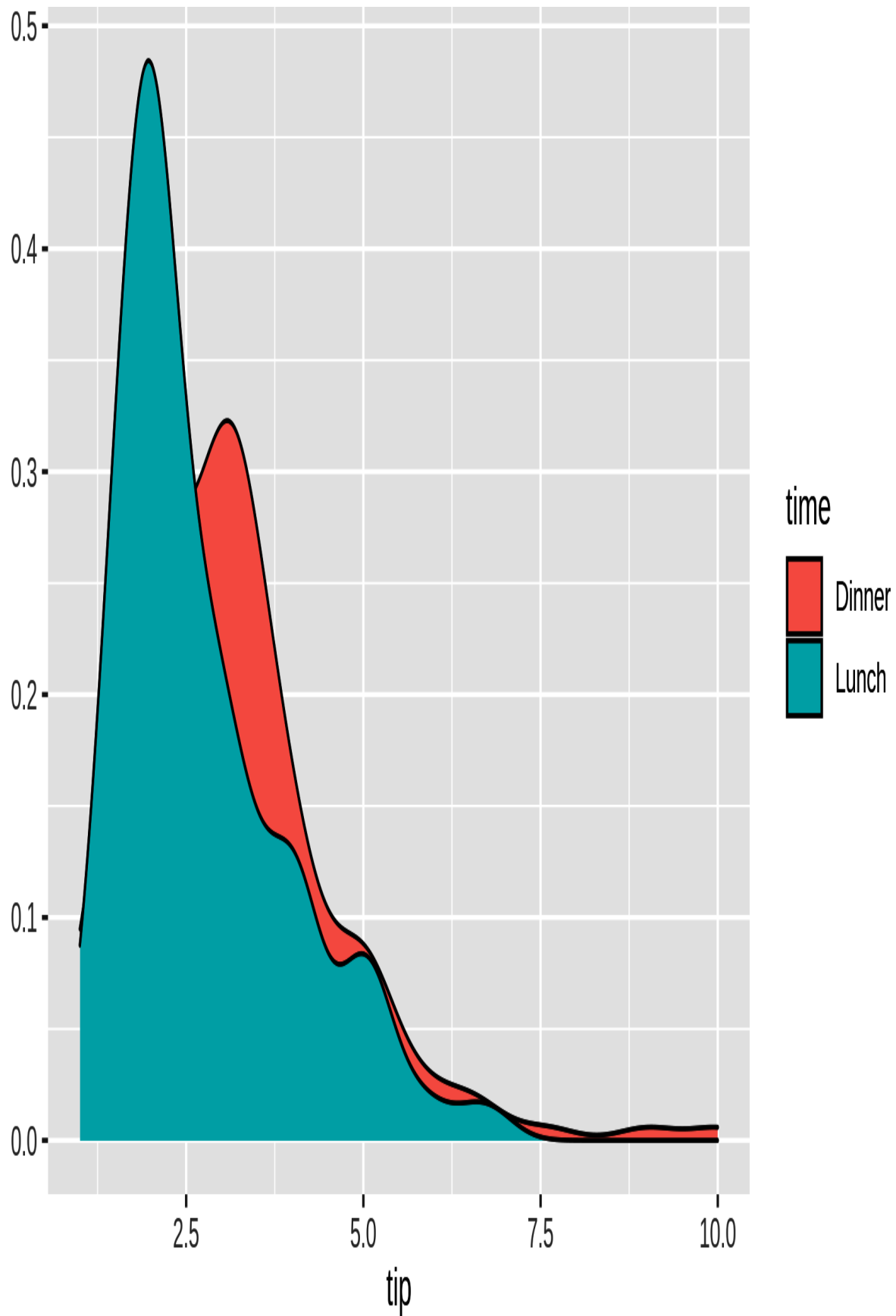
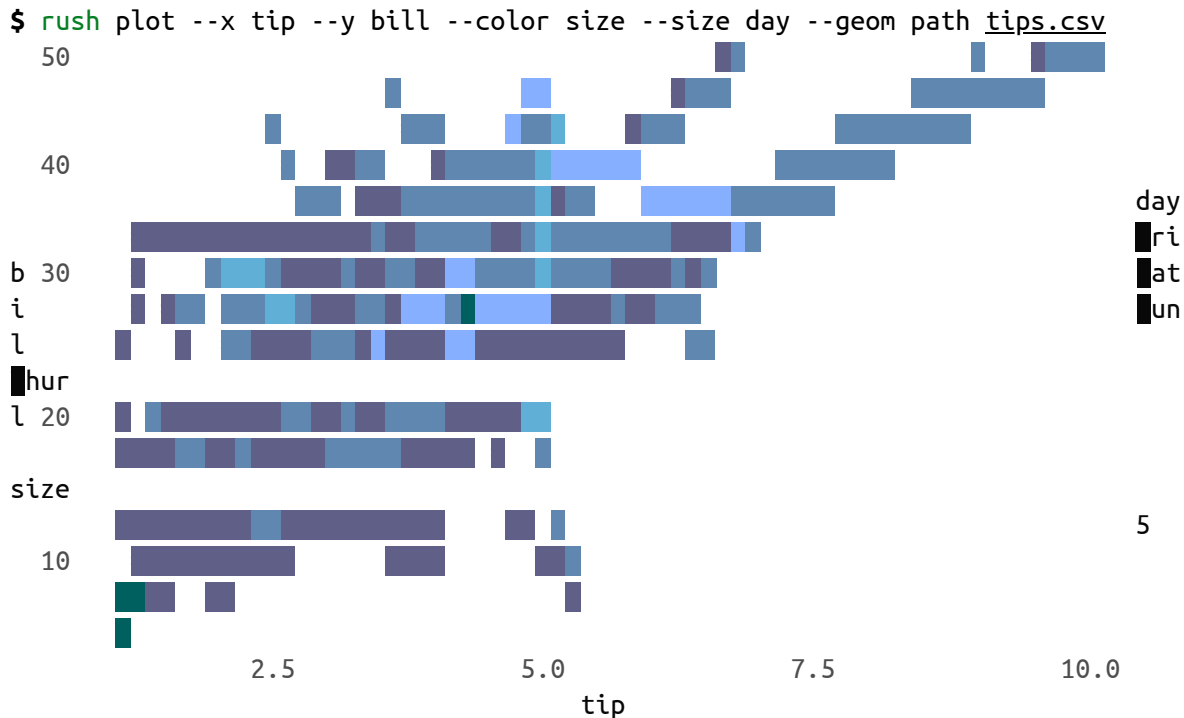


Figure 7-6. A density plot

Happy Little Accidents

You've already seen three types of visualizations. In ggplot2, these correspond to the functions `geom_bar`, `geom_histogram`, and `geom_density`. `geom` is short for *geometry* and dictates what is actually being plotted. This [cheat sheet for ggplot2](#) provides a good overview of the available geometry types. Which geometry types you can use depends on the columns that you specify (and their types). Not every combination makes sense. Take this line plot, for example:



This happy little accident becomes clearer in the visual representation in Figure 7-7:

```
$ rush plot --x tip --y bill --color size --size day --geom path tips.csv > plot  
-accident.png  
  
$ display plot-accident.png
```

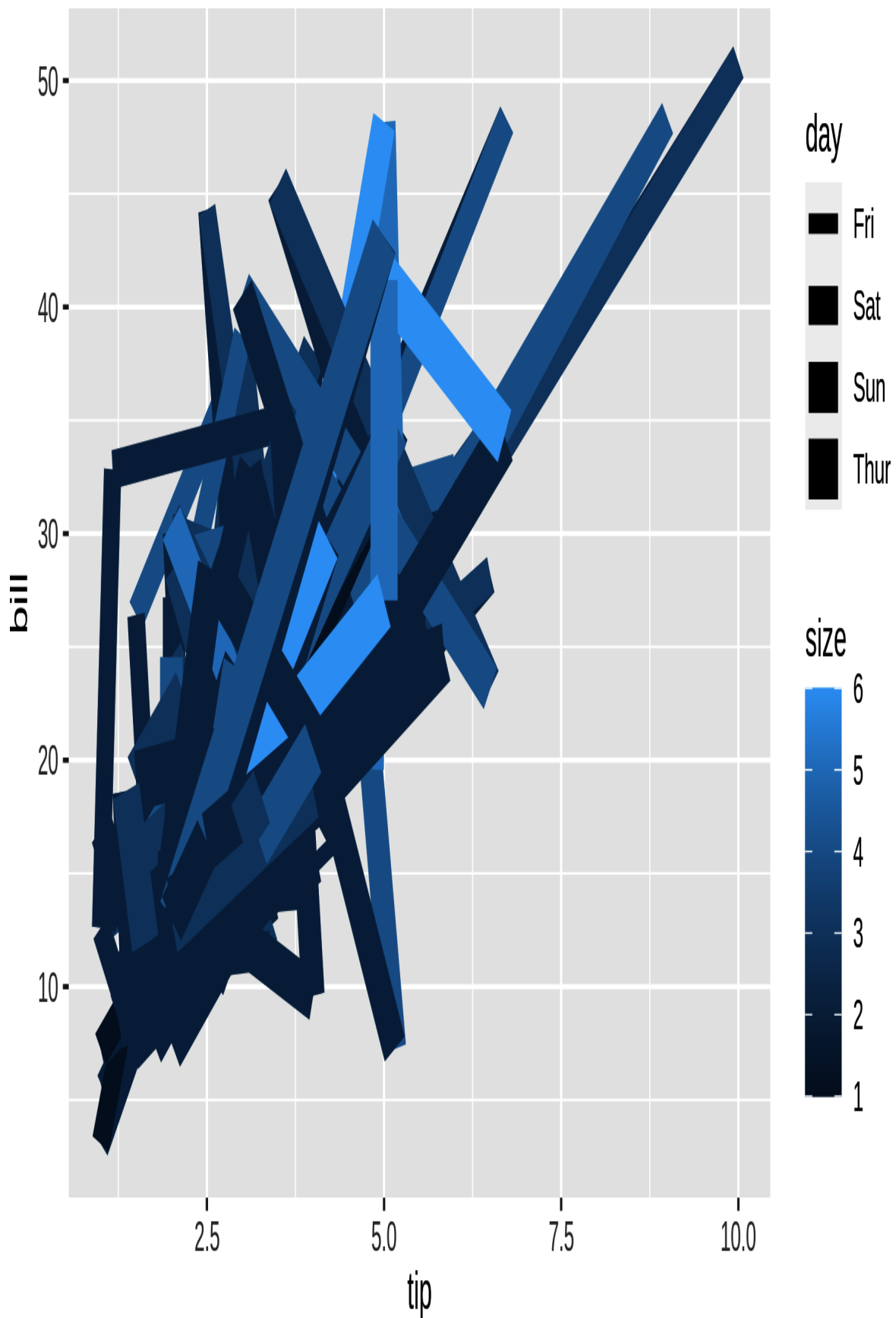


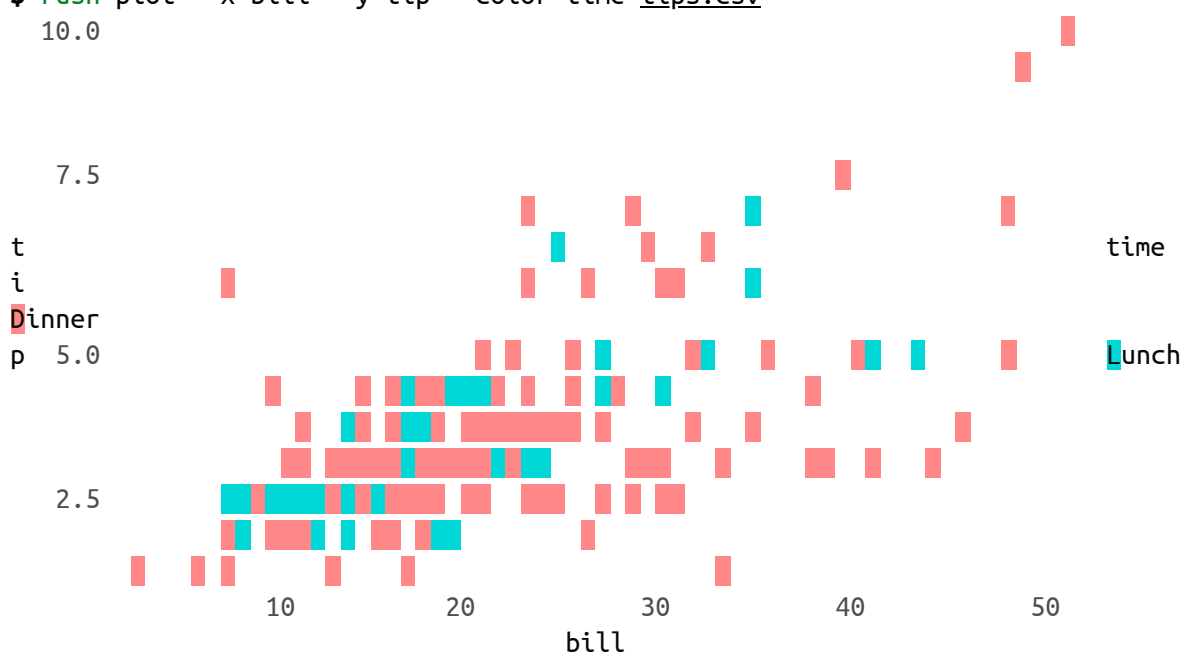
Figure 7-7. A happy little accident

The rows in `tips.csv` are independent observations, whereas drawing a line between the data points assumes that they are connected. It's better to visualize the relationship between the `tip` and the `bill` with a scatter plot.

Creating Scatter Plots

A scatter plot, where the geometry is a point, happens to be the default when specifying two continuous features:

```
$ rush plot --x bill --y tip --color time tips.csv  
10.0
```



Note that the color of each point is specified with the `--color` option (and not with the `--fill` option). See [Figure 7-8](#) for the visual representation:

```
$ rush plot --x bill --y tip --color time tips.csv > plot-scatter.png  
$ display plot-scatter.png
```

From this scatter plot we may conclude that there's a relationship between the amount of the bill and the tip. Perhaps it's useful to examine this data

from a higher level by creating trend lines.

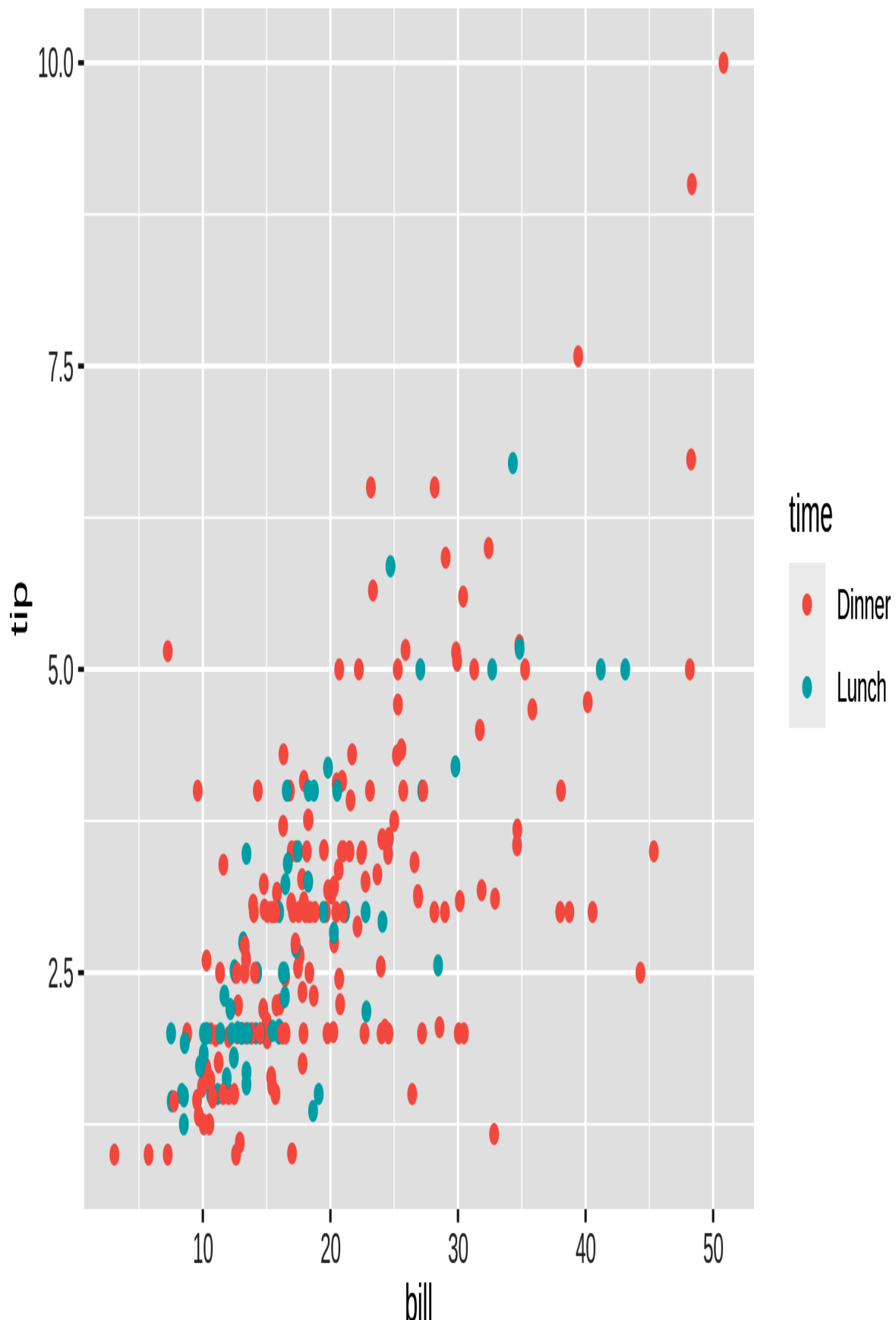
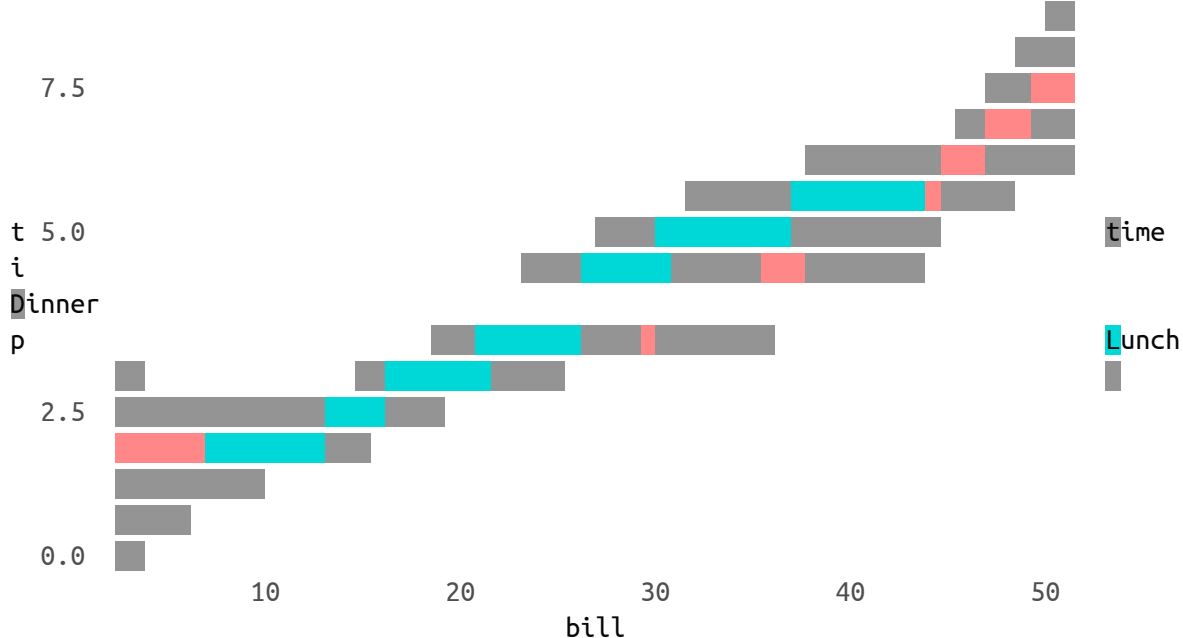


Figure 7-8. A scatter plot

Creating Trend Lines

If you override the default geometry with `smooth`, you can visualize trend lines. These are useful for seeing the bigger picture:

```
$ rush plot --x bill --y tip --color time --geom smooth tips.csv
```



`rush plot` cannot handle transparency, so a visual representation (see [Figure 7-9](#)) is much better in this case:

```
$ rush plot --x bill --y tip --color time --geom smooth tips.csv > plot-trend.png
$ display plot-trend.png
```

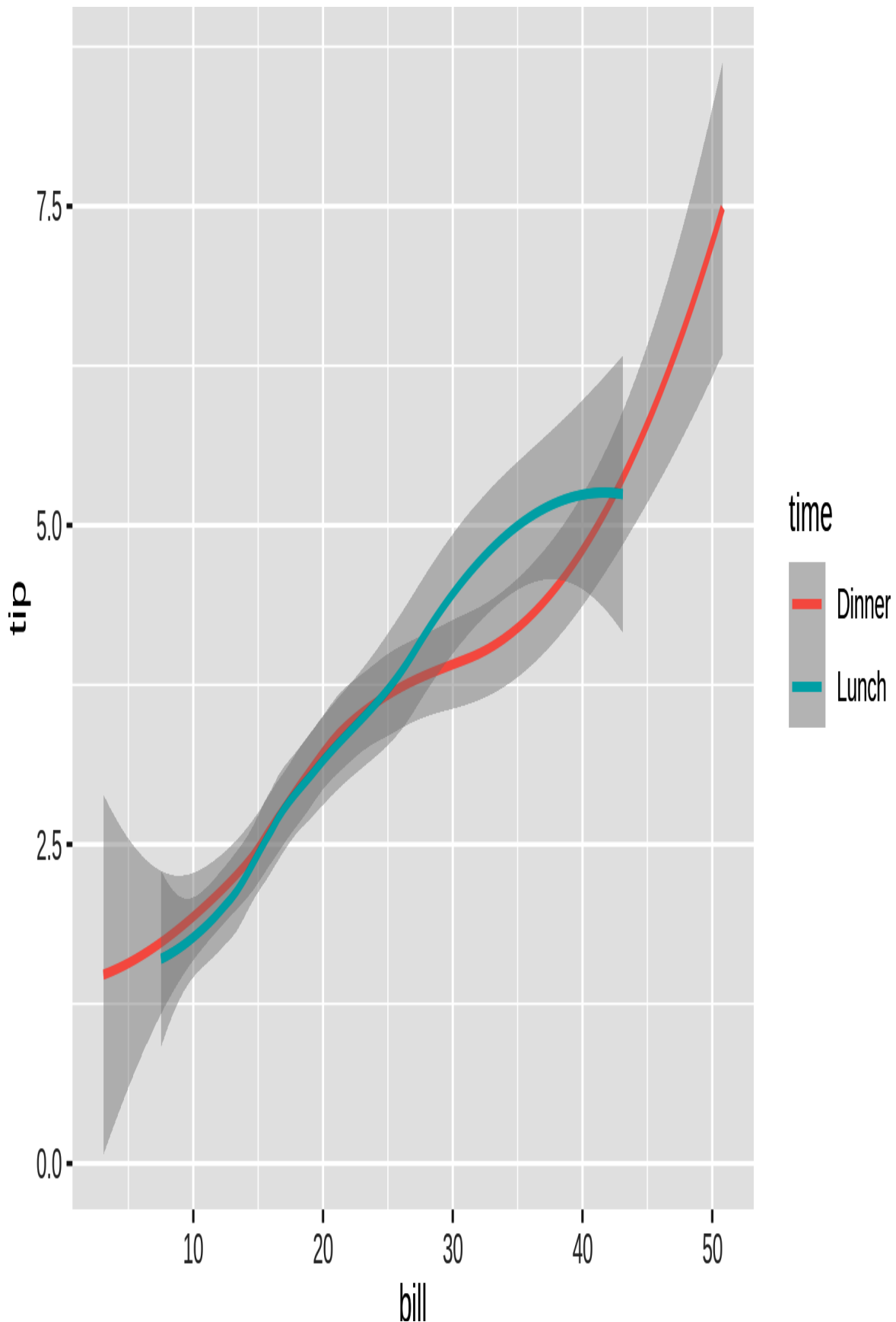


Figure 7-9. Trend lines

If you'd like to visualize the original points along with the trend lines, then you'll need to write ggplot2 code with `rush run` (see [Figure 7-10](#)):

```
$ rush run --library ggplot2 'ggplot(df, aes(x = bill, y = tip, color = time))  
+  
  geom_point() + geom_smooth()' tips.csv > plot-trend-points.png  
  
$ display plot-trend-points.png
```

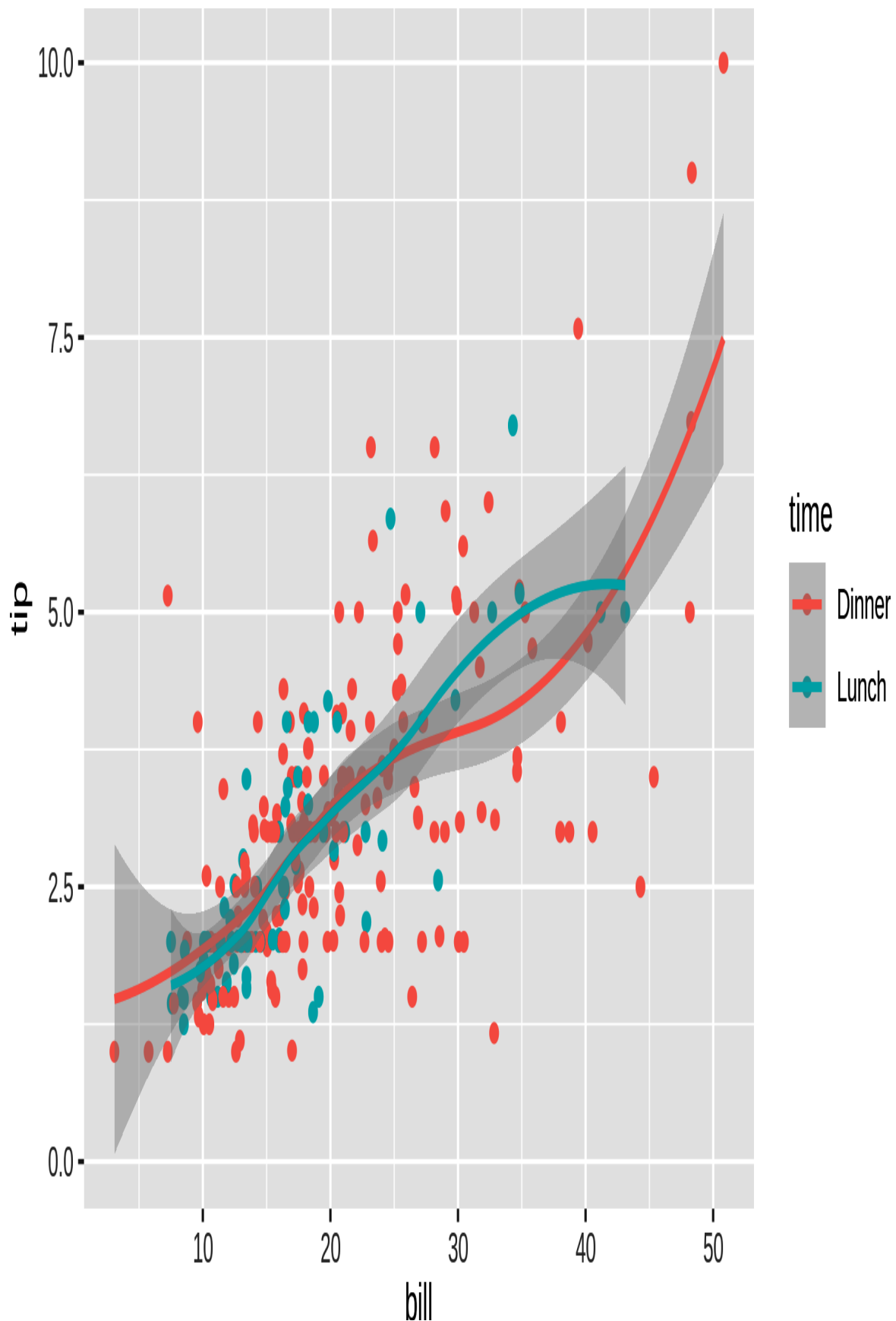
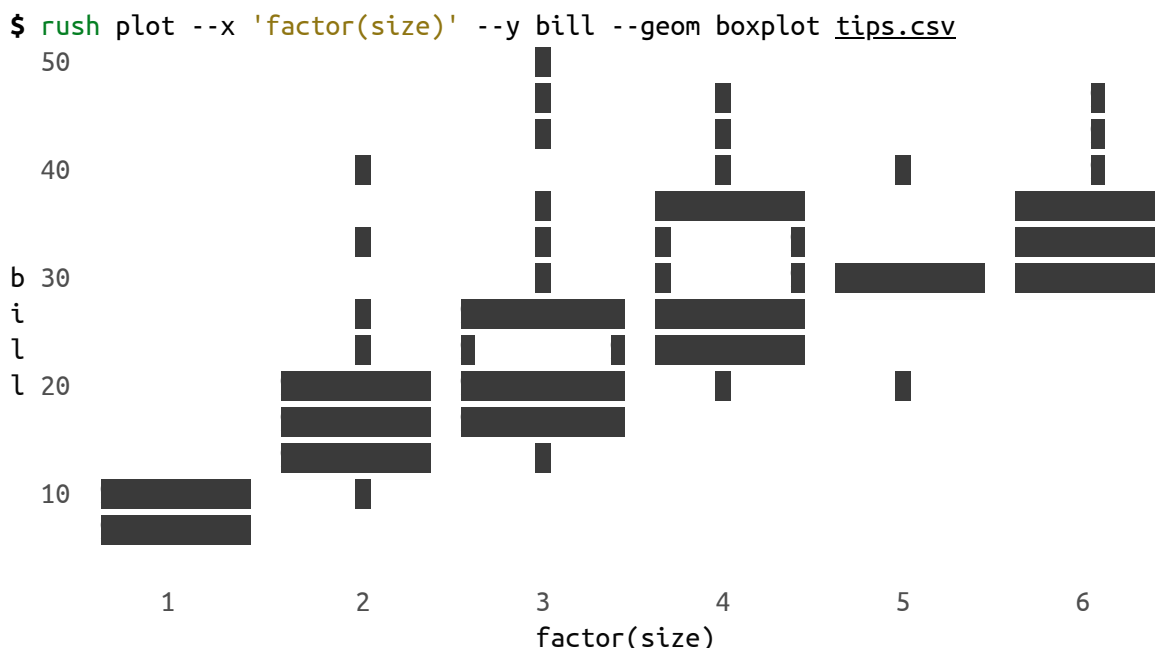


Figure 7-10. Trend lines and original points combined

Creating Box Plots

A box plot visualizes, for one or more features, a five-number summary: the minimum, the maximum, the sample median, and the first and third quartiles. In this case, we need to convert the *size* feature to a categorical one using the `factor()` function; otherwise, all values of the *bill* feature will be lumped together:



While the textual representation is not too bad, the visual one is much clearer (see Figure 7-11):

```
$ rush plot --x 'factor(size)' --y bill --geom boxplot tips.csv > plot-boxplot.png
```

```
$ display plot-boxplot.png
```

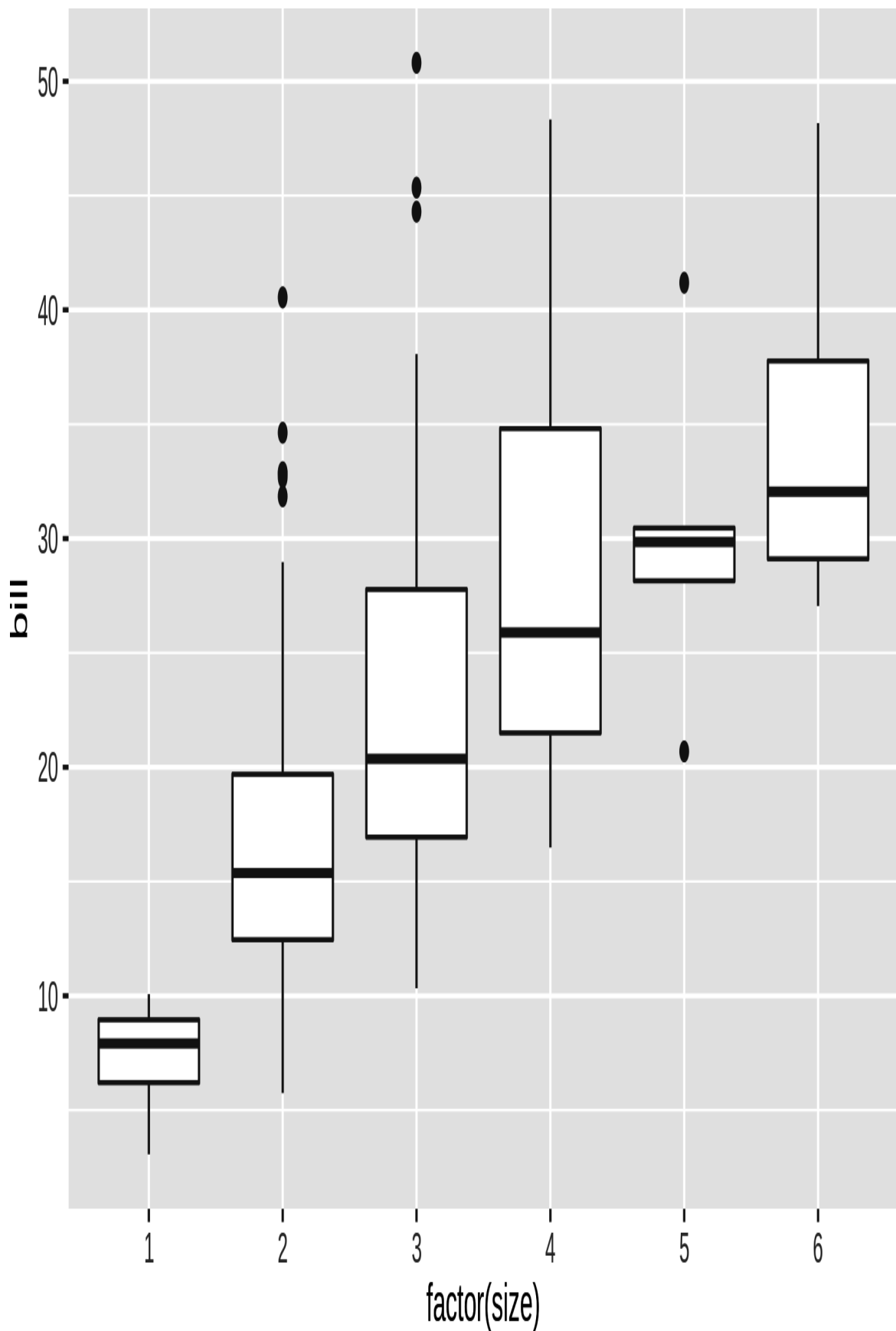
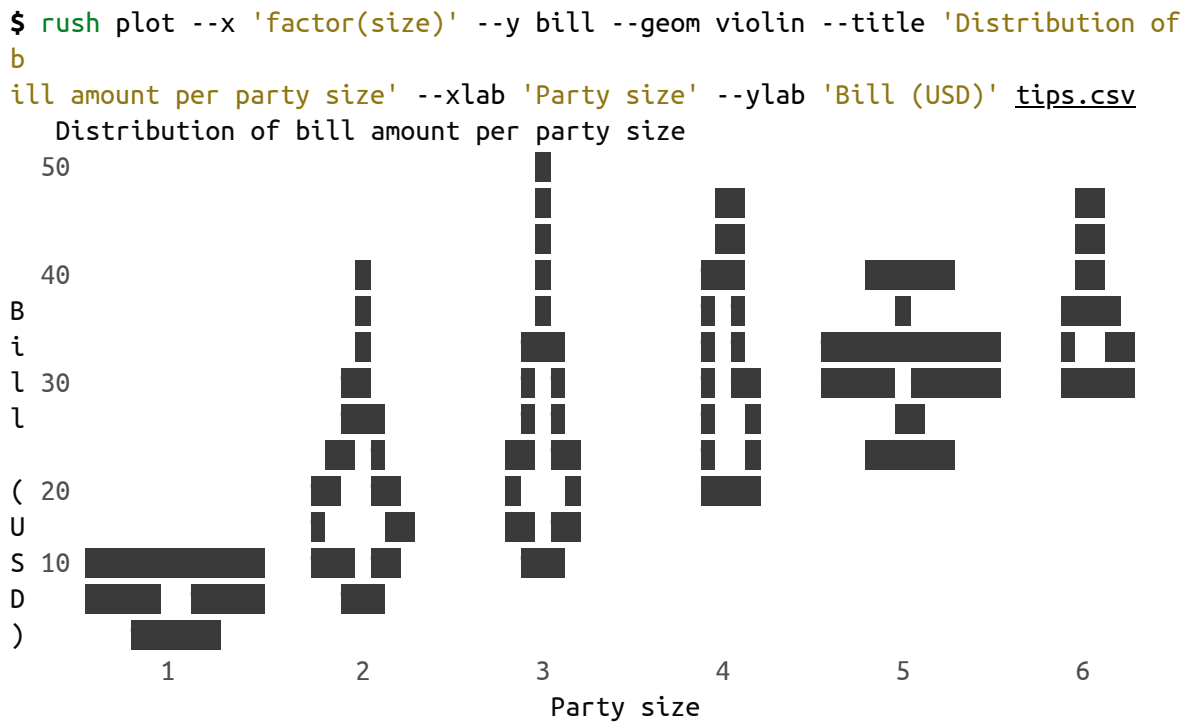


Figure 7-11. A box plot

Unsurprisingly, this box plot shows that, on average, a larger party size leads to a higher bill.

Adding Labels

The default labels are based on column names (or specifications). In the previous image, the label `factor(size)` should be improved. Using the `--xlab` and `--ylab` options, you can override the labels of the x- and y-axes. A title can be added with the `--title` option. Here's a violin plot (a mash-up of a box plot and a density plot) demonstrating this (see also [Figure 7-12](#)):



```
$ rush plot --x 'factor(size)' --y bill --geom violin --title 'Distribution of bill amount per party size' --xlab 'Party size' --ylab 'Bill (USD)' tips.csv > plot-labels.png
```

```
$ display plot-labels.png
```

Distribution of bill amount per party size

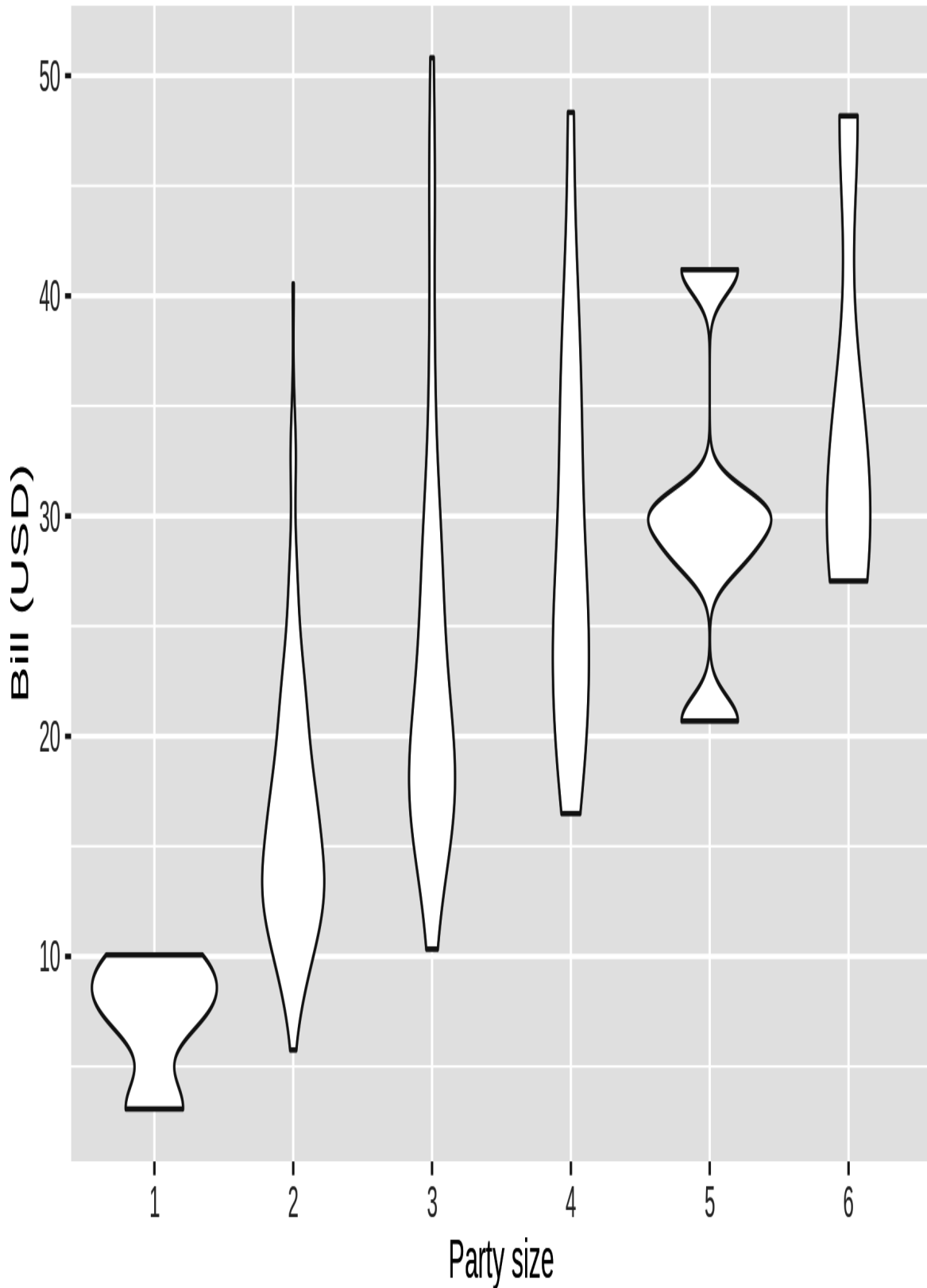


Figure 7-12. A violin plot with a title and labels

Annotating your visualization with proper labels and a title is especially useful if you want to share it with others (or your future self), as they will make it easier to understand what's being shown.

Going Beyond Basic Plots

Although `rush plot` is suitable for creating basic plots when you're exploring data, it certainly has its limitations. Sometimes you need more flexibility and sophisticated options such as multiple geometries, coordinate transformations, and theming. In that case, it might be worthwhile to learn more about the underlying package from which `rush plot` draws its capabilities, namely the `ggplot2` package for R. If you're more into Python than R, there's the [plotnine package](#), which is a reimplementation of `ggplot2` for Python.

Summary

In this chapter we've looked at various ways to explore your data. Both textual and graphical data visualizations have their pros and cons. The graphical ones are obviously of much higher quality but can be tricky to view at the command line. This is where textual visualizations come in handy. At least `rush`, thanks to R and `ggplot2`, has a consistent syntax for creating both types.

The next chapter is yet another intermezzo chapter in which I discuss how you can speed up your commands and pipelines. Feel free to read that chapter later if you can't wait to start modeling your data in [Chapter 9](#).

For Further Exploration

- A proper `ggplot2` tutorial is unfortunately beyond the scope of this book. If you want to get better at visualizing your data, I strongly recommend that you invest some time in understanding the power

and beauty of the grammar of graphics. Chapters 3 and 28 of the book *R for Data Science* by Hadley Wickham and Garrett Grolemund (O'Reilly) are an excellent resource.

- Speaking of Chapters 3 and 28, I translated those to Python using [plotnine](#) and [Pandas](#), in case you're more into Python than R.

1 Mark Nudelman, *less – Opposite of more*, version 551, 2019, <https://www.greenwoodsoftware.com/less>.

2 Christopher Groskopf, *csvstat – Print Descriptive Statistics for Each Column in a CSV File*, version 1.0.5, 2020, <https://csvkit.rtfd.org>.

3 The R Foundation for Statistical Computing, *R – a Language and Environment for Statistical Computing*, version 4.0.4, 2021, <https://www.r-project.org>.

4 Jeroen Janssens, *servewd – Serve the Current Working Directory Using a Simple HTTP Server*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.

Chapter 8. Parallel Pipelines

In the previous chapters, we've been dealing with commands and pipelines that take care of an entire task at once. In practice, however, you may find yourself facing a task that requires the same command or pipeline to run multiple times. For example, you may need to:

- Scrape hundreds of web pages
- Make dozens of API calls and transform their output
- Train a classifier for a range of parameter values
- Generate scatter plots for every pair of features in your dataset

In any of these examples, there's a certain form of repetition involved. With your favorite scripting or programming language, you could take care of this with a `for` loop or a `while` loop. On the command line, the first thing you might be inclined to do is to press the up arrow key to bring back the previous command, modify it if necessary, and press Enter to run the command again. This is fine to do two or three times, but imagine doing it *dozens* of times. Such an approach quickly becomes cumbersome, inefficient, and prone to errors. The good news is that you can write such loops on the command line as well. That's what this chapter is all about.

Sometimes, repeating a fast command again and again in succession (in a *serial* manner) is sufficient. When you have multiple cores (and perhaps even multiple machines), it would be nice to make use of those, especially when you're faced with a data-intensive task. Using multiple cores or machines may reduce the total running time significantly. In this chapter I will introduce a very powerful tool called `parallel`¹ that can take care of exactly this. It enables you to apply a command or pipeline for a range of arguments such as numbers, lines, and files. Plus, as the name implies, it allows you to run your commands in *parallel*.

Overview

This intermezzo chapter discusses several approaches to speeding up tasks that require commands and pipelines to be run many times. My main goal is to demonstrate to you the flexibility and power of `parallel`. Because this tool can be combined with any other tool discussed in this book, it will change the way you use the command line for data science for the better. In this chapter, you'll learn about:

- Running commands in serial to a range of numbers, lines, and files
- Breaking a large task into several smaller tasks
- Running pipelines in parallel
- Distributing pipelines to multiple machines

This chapter starts with the following files:

```
$ cd /data/ch08

$ l
total 20K
-rw-r--r-- 1 dst dst 126 Jun 29 14:32 emails.txt
-rw-r--r-- 1 dst dst 61 Jun 29 14:32 movies.txt
-rwxr-xr-x 1 dst dst 125 Jun 29 14:32 slow.sh*
-rw-r--r-- 1 dst dst 5.1K Jun 29 14:32 users.json
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Serial Processing

Before I dive into parallelization, I want to briefly discuss looping in a serial fashion. It's worthwhile to know how to do this because this functionality is always available, the syntax closely resembles looping in other programming languages, and it will really make you appreciate `parallel`.

From the examples provided in the introduction of this chapter, we can distill three types of items to loop over: numbers, lines, and files. These three types of items will be discussed in the next three subsections, respectively.

Looping Over Numbers

Imagine that you need to compute the square of every even integer between 0 and 100. There's a tool called `bc` that is a *basic calculator* you can pipe an equation to. The command to compute the square of 4 looks as follows:

```
$ echo "4^2" | bc  
16
```

For a one-off calculation, this will do. However, as mentioned in the introduction, you would have to be crazy to press the up arrow key, change the number, and press Enter 50 times! It would be better to let the shell do the hard work for you by using a `for` loop:

```
$ for i in {0..100..2}    ❶  
> do  
> echo "$i^2" | bc      ❷  
> done | trim  
0  
4  
16  
36  
64  
100  
144  
196  
256  
324  
... with 41 more lines
```

- ❶ The Z shell has a feature called *brace expansion*, which transforms `{0..100..2}` into a list separated by spaces: `0 2 4 ... 98 100`. The

variable `i` is assigned the value `0` in the first iteration, `1` in the second iteration, and so forth.

- ② The value of this variable can be used by prefixing it with a dollar sign (`$`). The shell will replace `$i` with its value before `echo` is executed.
Note that there can be more than one command between `do` and `done`.

Although the syntax may appear a bit odd compared to your favorite programming language, it's worth remembering this because it's always available in the shell. I'll introduce a better and more flexible way of repeating commands in a moment.

Looping Over Lines

The second type of item you can loop over is lines. These lines can come from either a file or standard input. This is a very generic approach because the lines can contain anything, including numbers, dates, and email addresses.

Imagine that you want to send an email to all your contacts. Let's first generate some fake users using the free [Random User Generator API](#):

```
$ curl -s "https://randomuser.me/api/1.2/?results=5&seed=dsatcl2e" > users.json
```

```
$ < users.json jq -r '.results[].email' > emails
```

```
$ bat emails
```

```
| File: emails
```

```
1 | selma.andersen@example.com
2 | kent.clark@example.com
3 | ditmar.niehaus@example.com
4 | benjamin.robinson@example.com
5 | paulo.muller@example.com
```

You can loop over the lines from *emails* with a `while` loop:

```
$ while read line
> do
> echo "Sending invitation to ${line}."      ②
> done < emails                            ③
Sending invitation to selma.andersen@example.com.
Sending invitation to kent.clark@example.com.
Sending invitation to ditmar.niehaus@example.com.
Sending invitation to benjamin.robinson@example.com.
Sending invitation to paulo.muller@example.com.
```

- ❶ In this case you need to use a `while` loop, because the Z shell does not know beforehand how many lines the input consists of.
- ❷ Although the curly braces around the `line` variable are not necessary in this case (since variable names cannot contain periods), including them is still good practice.
- ❸ This redirection can also be placed before `while`.

You can also provide input to a while loop interactively by specifying the special file standard input `/dev/stdin`. Press Ctrl-D when you are done:

```
$ while read line; do echo "You typed: ${line}."; done < /dev/stdin
one
You typed: one.
two
You typed: two.
three
You typed: three.
```

This method, however, has the disadvantage that, once you press Enter, the commands between `do` and `done` are run immediately for that line of input. There's no turning back.

Looping Over Files

In this section I discuss the third type of item that we often need to loop over: files.

To handle special characters, use *globbing* (i.e., pathname expansion) instead of `ls`:

```
$ for chapter in /data/*
> do
> echo "Processing Chapter ${chapter}."
> done
Processing Chapter /data/ch01.
Processing Chapter /data/ch02.
Processing Chapter /data/ch03.
Processing Chapter /data/ch04.
Processing Chapter /data/ch05.
Processing Chapter /data/ch06.
Processing Chapter /data/ch07.
Processing Chapter /data/ch08.
Processing Chapter /data/ch09.
Processing Chapter /data/ch10.
```

Just as with brace expansion, the expression `/data/*` is first expanded into a list by the Z shell before it's processed by the `for` loop. A more elaborate alternative to listing files is `find`,² which:

- Can traverse down directories
- Allows for elaborate searching on properties such as size, access time, and permissions
- Handles special characters such as spaces and newlines

For example, the following `find` invocation lists all files located under the directory `/data` that have `csv` as their extension and are smaller than 2 KB:

```
$ find /data -type f -name '*.csv' -size -2k
[data]/ch03/tmnt-basic.csv
[data]/ch03/tmnt-missing-newline.csv
[data]/ch03/tmnt-with-header.csv
[data]/ch05/names-comma.csv
[data]/ch05/irismeta.csv
```

```
/data/ch05/names.csv  
/data/ch07/datatypes.csv
```

Parallel Processing

Let's say that you have a very long running tool, such as the one shown here:

```
$ bat slow.sh  
-----  
|  
| File: slow.sh  
|-----  
|  
1 | #!/bin/bash  
2 | echo "Starting job $1" | ts ①  
3 | duration=$((1+RANDOM%5)) ②  
4 | sleep $duration ③  
5 | echo "Job $1 took ${duration} seconds" | ts  
-----  
|
```

- ① `ts` adds a timestamp.
- ② The magic variable `RANDOM` calls an internal Bash function that returns a pseudorandom integer between 0 and 32767. Taking the remainder of the division of that integer by 5 and adding 1 ensures that the *duration* is between 1 and 5.
- ③ `sleep` pauses execution for a given number of seconds.

This process probably doesn't take up all the available resources. And it so happens that you need to run this command a lot of times. For example, you need to download a whole sequence of files.

A naive way to parallelize is to run the commands in the background. Let's run `slow.sh` three times:

```

$ for i in {A..C}; do
> ./slow.sh $i & ❶
> done
[2] 162 ❷
[3] 163
[4] 164

$ Jun 29 14:32:36 Starting job A
Jun 29 14:32:36 Starting job B
Jun 29 14:32:36 Starting job C
Jun 29 14:32:37 Job B took 1 seconds

[3] - done      ./slow.sh $i
$ Jun 29 14:32:40 Job A took 4 seconds

[2] - done      ./slow.sh $i
$ Jun 29 14:32:41 Job C took 5 seconds

[4] + done      ./slow.sh $i
$
```

- ❶ The ampersand (&) sends the command to the background, allowing the for loop to continue immediately with the next iteration.
- ❷ This line shows the job number given by the Z shell and the process ID, which can be used for more fine-grained job control. This topic, while powerful, is beyond the scope of this book.

NOTE

Keep in mind that not everything can be parallelized. API calls may be limited to a certain number, and some commands can have only one instance.

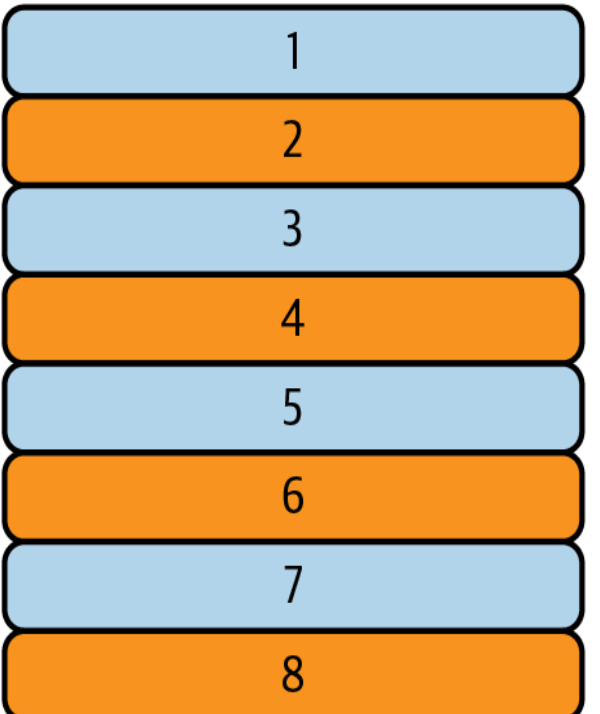
Figure 8-1 illustrates, on a conceptual level, the difference between serial processing, naive parallel processing, and parallel processing with GNU Parallel in terms of the number of concurrent processes and the total amount of time it takes to run everything.

There are two problems with the naive approach. First, there's no way to control how many processes you are running concurrently. If you start too many jobs at once, they could be competing for the same resources, such as CPU, memory, disk access, and network bandwidth. This could lead to taking longer to run everything. Second, it's difficult to tell which output belongs to which input. Let's look at a better approach.

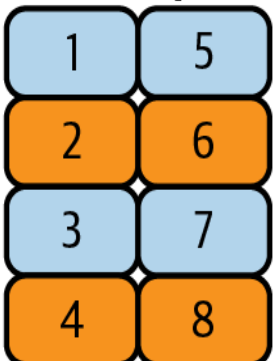
Serial processing



Naive parallel processing



Parallel processing with GNU parallel



Time →

Figure 8-1. Serial processing, naive parallel processing, and parallel processing with GNU Parallel

Introducing GNU Parallel

Allow me to introduce `parallel`, a command-line tool that allows you to parallelize and distribute commands and pipelines. The beauty of this tool is that existing tools can be used as they are; they do not need to be modified.

WARNING

Be aware that there are two command-line tools with the name `parallel`. If you're using the Docker image, then you already have the correct one installed. Otherwise, you can check that you have the correct one by running `parallel --version`. It should say "GNU parallel."

Before I delve into the details of `parallel`, here's a little teaser to show you how easy it is to replace the `for` loop from earlier:

```
$ seq 0 2 100 | parallel "echo {}^2 | bc" | trim
0
4
16
64
36
100
144
196
324
256
... with 41 more lines
```

This is `parallel` in its simplest form: the items to loop over are passed via standard input, and there aren't any arguments other than the command that `parallel` needs to run. See [Figure 8-2](#) for an illustration of how `parallel` concurrently distributes input among processes and collects their outputs.

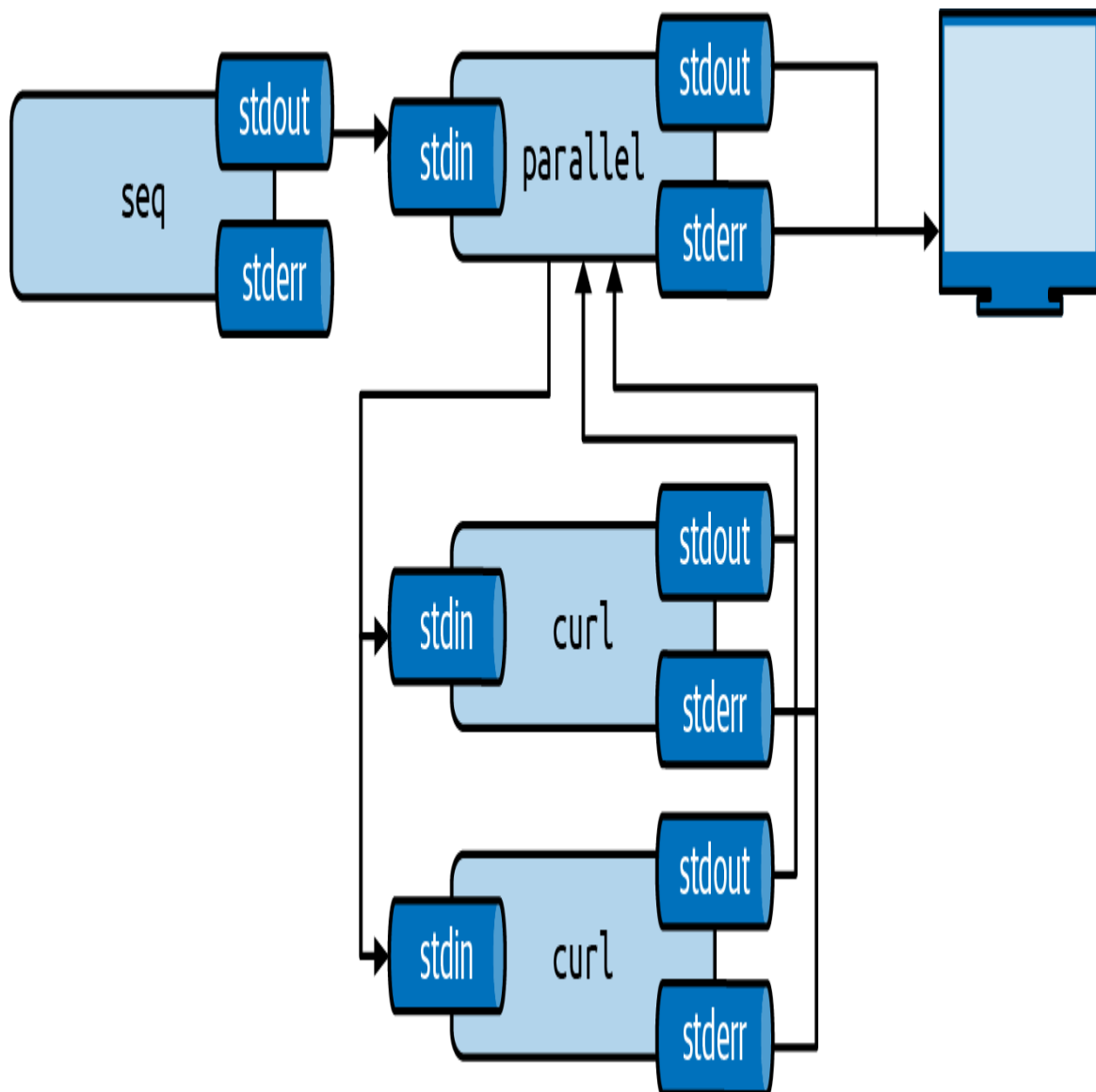


Figure 8-2. GNU Parallel concurrently distributes input among processes and collects their outputs

As you can see, it basically acts as a for loop. Here's another teaser, which replaces the for loop from the previous section:

```
$ parallel --jobs 2 ./slow.sh ::: {A..C}
Jun 29 14:32:44 Starting job B
Jun 29 14:32:47 Job B took 3 seconds
Jun 29 14:32:44 Starting job A
Jun 29 14:32:49 Job A took 5 seconds
Jun 29 14:32:47 Starting job C
Jun 29 14:32:51 Job C took 4 seconds
```

Here, using the `--jobs` option, I specify that `parallel` can run no more than two jobs concurrently. The arguments to `slow.sh` are specified as an argument instead of via standard input.

With a whopping 159 different options, `parallel` offers a lot of functionality. (Perhaps too much.) Luckily, you only need to know a handful to be effective. The manual page is quite informative, in case you need to use a less common option.

Specifying Input

The most important argument to `parallel` is the command or pipeline that you'd like to run for every input. The question is: where should the input item be inserted in the command line? If you don't specify anything, then the input item will be appended to the end of the pipeline:

```
$ seq 3 | parallel cowsay
```

```
< 1 >
---
\\  ^__^
 \\  (oo)\_____
   (__)\ )\/\
     ||----w |
     ||     |

< 2 >
---
\\  ^__^
 \\  (oo)\_____
   (__)\ )\/\
     ||----w |
     ||     |

< 3 >
---
\\  ^__^
 \\  (oo)\_____
   (__)\ )\/\
     ||----w |
     ||     |
```

The above is the same as running:

```
$ cowsay 1 > /dev/null ①
```

```
$ cowsay 2 > /dev/null
```

```
$ cowsay 3 > /dev/null
```

- ❶ Because the output is the same as before, I redirect it to */dev/null* to suppress it.

Although this often works, I advise you to be explicit about where the input item should be inserted in the command by using placeholders. In this case, because you want to use the entire input line (a number) at once, you need only one placeholder. You specify the placeholder—that is, where to put the input item—with a pair of curly braces ({}):

```
$ seq 3 | parallel cowsay {} > /dev/null
```

NOTE

There are other ways to provide input to `parallel`. I prefer piping the input (as I do throughout this chapter) because that's how most command-line tools are chained together into a pipeline. The other ways involve syntax that's not seen anywhere else. Having said that, they do enable additional functionality, such as iterating over all possible combinations of multiple lists, so be sure to read `parallel`'s manual page if you'd like to know more.

When the input items are filenames, some modifiers allow you to use only a portion of the filename. For example, with {}, only the basename of the filename will be used:

```
$ find /data/ch03 -type f | parallel echo '{#}\n\"{}\" has basename \"{}\"'  
①  
1) "/data/ch03/tmnt-basic.csv" has basename "tmnt-basic.csv"  
2) "/data/ch03/top2000.xlsx" has basename "top2000.xlsx"  
3) "/data/ch03/r-datasets.db" has basename "r-datasets.db"
```

```
4) "/data/ch03/tmnt-missing-newline.csv" has basename "tmnt-missing-newline.csv"  
5) "/data/ch03/tmnt-with-header.csv" has basename "tmnt-with-header.csv"  
6) "/data/ch03/logs.tar.gz" has basename "logs.tar.gz"
```

- ❶ Characters such as parentheses () and quotes ("") have a special meaning in the shell. To use them literally, you put a backslash (\) in front of them. This is called *escaping*.

If the input line has multiple parts separated by a delimiter, you can add numbers to the placeholders. For example:

```
$ < input.csv parallel --colsep , "mv {2} {1}" > /dev/null
```

Here, you can apply the same placeholder modifiers. It is also possible to reuse the same input item. If the input to **parallel** is a CSV file with a header, then you can use the column names as placeholders:

```
$ < input.csv parallel -C, --header : "invite {name} {email}"
```

TIP

If you ever start to wonder whether your placeholders are set up correctly, you can add the **--dryrun** option. Instead of actually executing the commands, **parallel** will print out all the commands exactly as if they have been executed.

Controlling the Number of Concurrent Jobs

By default, **parallel** runs one job per CPU core. You can control the number of jobs that will be run concurrently with the **--jobs** or **-j** option. Specifying a number (N) means that many jobs will be run concurrently. If you put a plus sign in front of the number, then **parallel** will run N jobs plus the number of CPU cores. If you put a minus sign in front of the number, then **parallel** will run $N - M$ jobs, where M is the number of CPU cores. You can also specify a percentage, where the default is 100% of

the number of CPU cores. The optimal number of jobs to run concurrently depends on the actual commands you are running:

```
$ seq 5 | parallel -j0 "echo Hi {}"  
Hi 1  
Hi 2  
Hi 3  
Hi 4  
Hi 5  
  
$ seq 5 | parallel -j200% "echo Hi {}"  
Hi 1  
Hi 2  
Hi 3  
Hi 4  
Hi 5
```

If you specify `-j1`, then the commands will be run in serial. Even though this approach doesn't do the name of the tool justice, it still has its uses—for example, when you need to access an API that allows only one connection at a time. If you specify `-j0`, then `parallel` will run as many jobs in parallel as possible. This can be compared to your `for` loop with the ampersand. Doing this is not advised.

Logging and Output

To save the output of each command, you might be tempted to do the following:

```
$ seq 5 | parallel "echo \"Hi {}\" > hi-{}.txt"
```

This will save the output into individual files. Or if you want to save everything into one big file, you could do the following:

```
$ seq 5 | parallel "echo Hi {}" >> one-big-file.txt
```

However, `parallel` offers the `--results` option, which stores the output in separate files. For each job, `parallel` creates three files: `seq`, which

holds the job number; *stdout*, which contains the output produced by the job; and *stderr*, which contains any errors produced by the job. These three files are placed in subdirectories based on the input values.

`parallel` still prints all the output, which is redundant in this case. You can redirect both the standard input and standard output to `/dev/null` as follows:

```
$ seq 10 | parallel --results outdir "curl  
'https://anapioficeandfire.com/api/ch  
aracters/{}' | jq -r '.aliases[0]'" 2>/dev/null 1>&2  
  
$ tree outdir | trim  
outdir  
└── 1  
    ├── 1  
    │   ├── seq  
    │   ├── stderr  
    │   └── stdout  
    └── 10  
        ├── seq  
        ├── stderr  
        └── stdout  
... with 34 more lines
```

See [Figure 8-3](#) for a pictorial overview of how the `--results` option works.

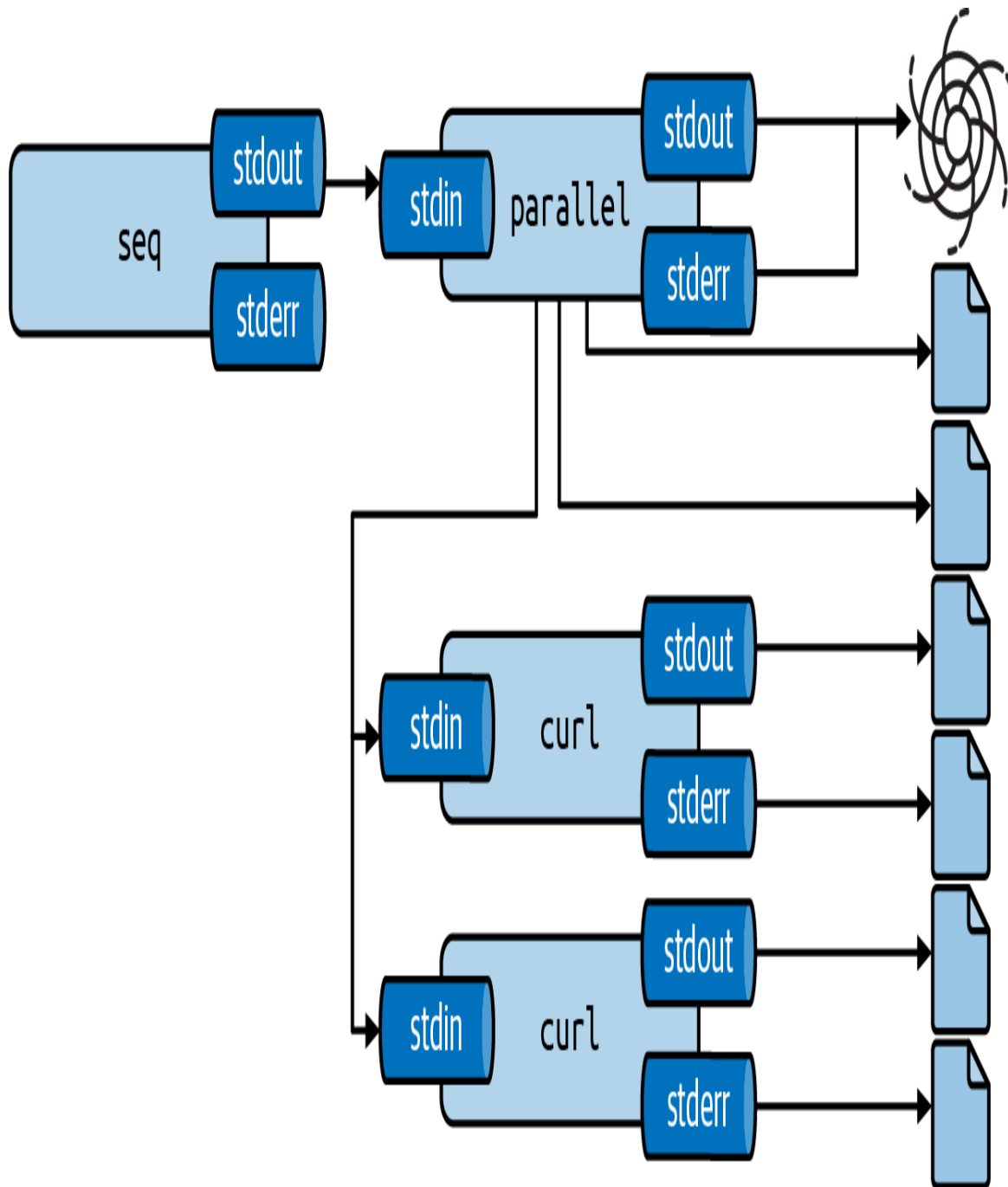


Figure 8-3. GNU Parallel stores output in separate files with the `--results` option

When you're running multiple jobs in parallel, the order in which the jobs are run may not correspond to the order of the input. The output of jobs is therefore also mixed up. To keep the same order, specify the `--keep-order` or `-k` option.

Sometimes it's useful to record which input generated which output. `parallel` allows you to *tag* the output with the `--tag` option, which prepends each line with the input item:

```
$ seq 5 | parallel --tag "echo 'sqrt({})' | bc -l"
1      1
2      1.41421356237309504880
3      1.73205080756887729352
4      2.000000000000000000000000
5      2.23606797749978969640

$ parallel --tag --keep-order "echo '{1}*{2}' | bc -l" :::: 3 4 :::: 5 6 7
3 5      15
3 6      18
3 7      21
4 5      20
4 6      24
4 7      28
```

Creating Parallel Tools

The `bc` tool, which I used in the beginning of this chapter, is not parallel by itself. However, you can parallelize it using `parallel`. The Docker image contains a tool called `pbc`.³ Its code is shown here:

```
$ bat $(which pbc)
-----
| File: /usr/bin/dsutils/pbc
-----
|
1  |#!/bin/bash
2  |# pbc: parallel bc. First column of input CSV is mapped to {1},
second
    | to {2}, and so forth.
3  |
4  |# Example usage: paste -d, <(seq 100) <(seq 100 -1 1) | ./pbc
'sqrt({1}
    | *{2})'
5  |
6  |# Dependency: GNU parallel
7  |
8  |# Author: http://jeroenjanssens.com
9  |
```

```
10 | parallel -C, -k -j100% "echo '$1' | bc -l"
```

This tool allows us to simplify the code used in the beginning of the chapter too. And it can process comma-separated values simultaneously:

```
$ seq 100 | pbc '{1}^2' | trim  
1  
4  
9  
16  
25  
36  
49  
64  
81  
100  
... with 90 more lines  
  
$ paste -d, <(seq 4) <(seq 4) <(seq 4) | pbc 'sqrt({1}+{2})^{3}'  
1.41421356237309504880  
4.000000000000000000000000  
14.69693845669906858905  
63.999999999999999999999969
```

Distributed Processing

Sometimes you need more power than your local machine, even with all its cores, can offer. Luckily, `parallel` can also leverage the power of remote machines, which really allows you to speed up your pipeline.

What's great is that `parallel` doesn't have to be installed on the remote machine. All that's required is that you're able to connect to the remote machine with the *Secure Shell* protocol (or SSH), which is also what `parallel` uses to distribute your pipeline. (Having `parallel` installed is helpful because it can then determine how many cores to employ on each remote machine; I'll talk more about this later.)

First, we're going to obtain a list of running Amazon Web Services' Elastic Compute Cloud (AWS EC2) instances. Don't worry if you don't have any

remote machines; you can replace any occurrence of `--slf hostnames`, which tells `parallel` which remote machines to use, with `--sshlogin :.`. This way, you can still follow along with the examples in this section.

Once you know which remote machines to take over, there are three flavors of distributed processing to consider:

- Running ordinary commands on remote machines
- Distributing local data directly among remote machines
- Sending files to remote machines, processing them, and retrieving the results

Get List of Running AWS EC2 Instances

In this section, we're creating a file named `hostnames` that will contain one hostname of a remote machine per line. I'm using Amazon Web Services (AWS) as an example. I assume that you have an AWS account and that you know how to launch instances. If you're using a different cloud computing service (such as Google Cloud Platform or Microsoft Azure), or if you have your own servers, please make sure that you create a `hostnames` file yourself before continuing to the next section.

You can obtain a list of running AWS EC2 instances using `aws`,⁴ the command-line interface to the AWS API. With `aws`, you can do almost everything you can do with the online AWS Management Console.

The command `aws ec2 describe-instances` returns a lot of information about all your EC2 instances in JSON format (see [the online documentation](#) for more information). You can extract the relevant fields using `jq`:

```
$ aws ec2 describe-instances | jq '.Reservations[].Instances[] | {public_dns: .PublicIpAddress, state: .State.Name}'
```

The possible states of an EC2 instance are `pending`, `running`, `shutting-down`, `terminated`, `stopping`, and `stopped`. Because you can only

distribute your pipeline to running instances, you filter out the nonrunning instances as follows:

```
> aws ec2 describe-instances | jq -r '.Reservations[].Instances[] |  
select(.Stat  
e.Name=="running") | .PublicDnsName' | tee hostnames  
ec2-54-88-122-140.compute-1.amazonaws.com  
ec2-54-88-89-208.compute-1.amazonaws.com
```

(Without the `-r` or `--raw-output` option, the hostnames would have been surrounded by double quotes.) The output is saved to *hostnames*, so that I can pass this to `parallel` later.

As mentioned, `parallel` employs `ssh5` to connect to the remote machines. If you want to connect to your EC2 instances without typing the credentials every time, you can add something like the following text to the file `~/.ssh/config`:

```
$ bat ~/.ssh/config
```

```
-----  
|  
| File: /home/dst/.ssh/config  
|  
|-----  
| 1 | Host *.amazonaws.com  
| 2 |     IdentityFile ~/.ssh/MyKeyFile.pem  
| 3 |     User ubuntu  
|-----  
|  
|-----
```

Depending on which distribution you're running, your username may be different than `ubuntu`.

Running Commands on Remote Machines

The first flavor of distributed processing is to run ordinary commands on remote machines. Let's first double-check that `parallel` is working by running the tool `hostname6` on each EC2 instance:

```
$ parallel --nonall --sshloginfile hostnames hostname
ip-172-31-23-204
ip-172-31-23-205
```

Here, the `--sshloginfile` or `--slf` option is used to refer to the file `hostnames`. The `--nonall` option instructs `parallel` to execute the same command on every remote machine in the `hostnames` file without using any parameters. Remember, if you don't have any remote machines to utilize, you can replace `--slf hostnames` with `--sshlogin :` so that the command is run on your local machine:

```
$ parallel --nonall --sshlogin : hostname
data-science-toolbox
```

Running the same command on every remote machine once requires only one core per machine. If you wanted to distribute the list of arguments passed in to `parallel`, then it could potentially use more than one core. If it's not specified explicitly, `parallel` will try to determine the number of cores:

```
$ seq 2 | parallel --slf hostnames echo 2>&1
bash: parallel: command not found
parallel: Warning: Could not figure out number of cpus on ec2-54-88-122-
140.comp
ute-1.amazonaws.com (). Using 1.
1
2
```

In this case, I have `parallel` installed on one of the two remote machines. I'm getting a warning message indicating that `parallel` is not found on one of them. As a result, `parallel` cannot determine the number of cores and will default to using one core. When you receive this warning message, you can do one of the following four things:

- Don't worry and be happy with using one core per machine.
- Specify the number of jobs for each machine via the `--jobs` or `-j` option.

- Specify the number of cores to use per machine by putting, for example, `2/` (if you want two cores), in front of each hostname in the `hostnames` file.
- Install `parallel` using a package manager—for example, if the remote machines all run Ubuntu:

```
$ parallel --nonall --slf hostnames "sudo apt-get install -y parallel"
```

Distributing Local Data Among Remote Machines

The second flavor of distributed processing is to distribute local data directly among remote machines. Imagine that you have one very large dataset that you want to process using multiple remote machines. For simplicity's sake, let's sum all integers from 1 to 1,000. First, let's double-check that your input is actually being distributed by printing the hostname of the remote machine and the length of the input it received using `wc`:

```
$ seq 1000 | parallel -N100 --pipe --slf hostnames "(hostname; wc -l) | paste -s d:"  
ip-172-31-23-204:100  
ip-172-31-23-205:100  
ip-172-31-23-205:100  
ip-172-31-23-204:100  
ip-172-31-23-205:100  
ip-172-31-23-204:100  
ip-172-31-23-205:100  
ip-172-31-23-204:100  
ip-172-31-23-205:100  
ip-172-31-23-204:100
```

Excellent. You can see that your 1,000 numbers get distributed evenly in subsets of 100 (as specified by `-N100`). Now you're ready to sum all those numbers:

```
$ seq 1000 | parallel -N100 --pipe --slf hostnames "paste -sd+ | bc" | paste -sd  
500500
```

Here, you immediately also sum the 10 sums you get back from the remote machines. Let's check that the answer is correct by doing the same calculation without `parallel`:

```
$ seq 1000 | paste -sd+ | bc  
500500
```

Good, that works. If you have a larger pipeline that you want to execute on the remote machines, you can also put it in a separate script and upload it with `parallel`. I'll demonstrate this by creating a very simple command-line tool called `add`:

```
$ echo '#!/usr/bin/env bash' > add  
$ echo 'paste -sd+ | bc' >> add  
  
$ bat add  
---  
| File: add  
---  
|  
| 1 | #!/usr/bin/env bash  
| 2 | paste -sd+ | bc  
---  
  
$ chmod u+x add  
  
$ seq 1000 | ./add  
500500
```

Using the `--basefile` option, `parallel` first uploads the file `add` to all remote machines before running the jobs:

```
$ seq 1000 |  
> parallel -N100 --basefile add --pipe --slf hostnames './add' |  
> ./add  
500500
```

Summing 1,000 numbers is of course only a toy example. And it would've been much faster to do this locally. Still, I hope it's clear from this that `parallel` can be incredibly powerful.

Processing Files on Remote Machines

The third flavor of distributed processing is to send files to remote machines, process them, and retrieve the results. Imagine that you want to count how often each borough of New York City receives service calls on 311. You don't have that data on your local machine yet, so let's first obtain it from the free [NYC Open Data API](#):

```
$ seq 0 100 900 | parallel "curl -sL  
'http://data.cityofnewyork.us/resource/erm  
2-nwe9.json?\\$limit=100&\\$offset={} | jq -c '.[]' | gzip > nyc-{}.json.gz"
```

You now have 10 files containing compressed JSON data:

```
$ ls nyc*.json.gz  
-rw-r--r-- 1 dst dst 14K Jun 29 14:34 nyc-10.json.gz  
-rw-r--r-- 1 dst dst 13K Jun 29 14:33 nyc-1.json.gz  
-rw-r--r-- 1 dst dst 13K Jun 29 14:33 nyc-2.json.gz  
-rw-r--r-- 1 dst dst 14K Jun 29 14:33 nyc-3.json.gz  
-rw-r--r-- 1 dst dst 14K Jun 29 14:33 nyc-4.json.gz  
-rw-r--r-- 1 dst dst 13K Jun 29 14:33 nyc-5.json.gz  
-rw-r--r-- 1 dst dst 14K Jun 29 14:34 nyc-6.json.gz  
-rw-r--r-- 1 dst dst 13K Jun 29 14:33 nyc-7.json.gz  
-rw-r--r-- 1 dst dst 14K Jun 29 14:34 nyc-8.json.gz  
-rw-r--r-- 1 dst dst 14K Jun 29 14:34 nyc-9.json.gz
```

Note that `jq -c '.[]'` is used to flatten the array of JSON objects so that there's one object per line, with a total of 100 lines per file. Using `zcat`,⁷ you can directly print the contents of a compressed file:

```
$ zcat nyc-1.json.gz | trim  
{"unique_key": "51022229", "created_date": "2021-06-  
28T02:00:31.000", "agency": "NYP...  
{"unique_key": "51026918", "created_date": "2021-06-  
28T02:00:10.000", "agency": "NYP...  
{"unique_key": "51024237", "created_date": "2021-06-
```

```

28T02:00:04.000", "agency": "NYP...
{"unique_key": "51025570", "created_date": "2021-06-
28T01:59:56.000", "agency": "NYP...
{"unique_key": "51021715", "created_date": "2021-06-
28T01:59:54.000", "agency": "NYP...
{"unique_key": "51029212", "created_date": "2021-06-
28T01:59:46.000", "agency": "NYP...
{"unique_key": "51026777", "created_date": "2021-06-
28T01:59:42.000", "agency": "NYP...
{"unique_key": "51028611", "created_date": "2021-06-
28T01:59:32.000", "agency": "NYP...
{"unique_key": "51019729", "created_date": "2021-06-
28T01:59:28.000", "agency": "NYP...
{"unique_key": "51028159", "created_date": "2021-06-
28T01:59:17.000", "agency": "NYP...
... with 90 more lines

```

Let's see what one line of JSON looks like using `head`:

```

$ zcat nyc-1.json.gz | head -n 1
{"unique_key": "51022229", "created_date": "2021-06-
28T02:00:31.000", "agency": "NYPD
", "agency_name": "New York City Police Department", "complaint_type": "Blocked
Driv
eway", "descriptor": "No
Access", "location_type": "Street/Sidewalk", "incident_zip": "10031", "incident_address": "791 ST NICHOLAS AVENUE", "street_name": "ST NICHOLAS
A
VENUE", "cross_street_1": "WEST 149 STREET", "cross_street_2": "WEST 150
STREET", "
intersection_street_1": "WEST 149 STREET", "intersection_street_2": "WEST 150
STR
EET", "city": "NEW YORK", "landmark": "ST NICHOLAS AVENUE", "status": "In
Progress", "c
ommunity_board": "09
MANHATTAN", "bbl": "1020640033", "borough": "MANHATTAN", "x_coord
inate_state_plane": "1000047", "y_coordinate_state_plane": "240589", "open_data_ch
an
nel_type": "MOBILE", "park_facility_name": "Unspecified", "park_borough": "MANHATTA
N"
, "latitude": "40.827022608423036", "longitude": "-73.94292011775158", "location": {"l
atitude": "40.827022608423036", "longitude": "-73.94292011775158", "human_address": ":"}
{\"address\": \"\", \"city\": \"\", \"state\": \"\", \"zip\": \"\"}, :@compute

```

```
d_region_efsh_h5xi":"12428",":@computed_region_f5dn_yrer":"37",":@computed_region_i
io
n_yeji_bk3q":"4",":@computed_region_92fq_4b7q":"36",":@computed_region_sbqj_en
ih
":"19"}
```

If you wanted to get the total number of service calls per borough on your local machine, you would run the following command:

```
$ zcat nyc*.json.gz | ①
> jq -r '.borough' | ②
> tr '[A-Z]' '[a-z]_' | ③
> sort | uniq -c | sort -nr | ④
> awk '{print $2","$1}' | ⑤
> header -a borough,count | ⑥
> csvlook
| borough      | count |
|-----+-----|
| bronx        | 349   |
| manhattan    | 283   |
| brooklyn     | 218   |
| queens        | 137   |
| staten_island | 13    |
```

- ❶ Expand all compressed files using `zcat`.
- ❷ For each call, extract the name of the borough using `jq`.
- ❸ Convert borough names to lowercase and replace spaces with underscores (because `awk` splits on whitespace by default).
- ❹ Count the occurrences of each borough using `sort` and `uniq`.
- ❺ Reverse the two columns and delimit them by comma using `awk`.
- ❻ Add a header using `header`.

Imagine for a moment that your own machine is so slow that you simply cannot perform this pipeline locally. You can use `parallel` to distribute the local files among the remote machines, let them do the processing, and retrieve the results:

```
$ ls *.json.gz | ❶
> parallel -v --basefile jq \ ❷
> --trc {.}.csv \ ❸
> --slf hostnames \ ❹
> "zcat {} | ./jq -r '.borough' | tr '[A-Z]' '[a-z]_' | sort | uniq -c | awk
'{
print \"$2\", \"$1\" > {.}.csv" ❺
```

- ❶ Print the list of files and pipe it into `parallel`.
- ❷ Transmit the `jq` binary to each remote machine. Luckily, `jq` has no dependencies. This file will be removed from the remote machines afterward because I specified the `--trc` option (which implies the `--cleanup` option). Note that the pipeline uses `./jq` instead of just `jq`. That's because the pipeline needs to use the version that was uploaded and not the version that may or may not be on the search path.
- ❸ The command-line argument `--trc {.}.csv` is short for `--transfer --return {.}.csv --cleanup`. (The replacement string `{.}` gets replaced with the input filename without the last extension.) Here, this means that the JSON file gets transferred to the remote machine, the CSV file gets returned to the local machine, and both files will be removed from the remote machine after each job.
- ❹ Specify a list of hostnames. Remember, if you want to try this out locally, you can specify `--sshlogin :` instead of `--slf hostnames`.
- ❺ Note the escaping in the `awk` expression. Quoting can sometimes be tricky. Here, the dollar signs and the double quotes are escaped. If

quoting ever gets too confusing, remember that you put the pipeline into a separate command-line tool, just as I did with `add`.

If you were to run `ls` on one of the remote machines during this process, you would see that `parallel` indeed transfers (and cleans up) the binary `jq`, the JSON files, and the CSV files:

```
$ ssh $(head -n 1 hostnames) ls
```

Each CSV file looks something like this:

```
> cat nyc-1.json.csv
bronx,3
brooklyn,5
manhattan,24
queens,3
staten_island,2
```

You can sum the counts in each CSV file by using `rush` and the tidyverse:

```
$ cat nyc*csv | header -a borough,count |
> rush run -t 'group_by(df, borough) %>% summarize(count = sum(count))' - |
> csvsort -rc count | csvlook
| borough      | count |
|-----|
| bronx        | 349 |
| manhattan    | 283 |
| brooklyn     | 218 |
| queens        | 137 |
| staten_island | 13 |
```

Or if you prefer to use SQL to aggregate results, you can use `csvsql`, as discussed in [Chapter 5](#):

```
$ cat nyc*csv | header -a borough,count |
> csvsql --query 'SELECT borough, SUM(count) AS count FROM stdin GROUP BY
  borough ORDER BY count DESC' |
> csvlook
| borough      | count |
|-----|
```

bronx	349
manhattan	283
brooklyn	218
queens	137
staten_island	13

Summary

As a data scientist, you work with data—occasionally a lot of data. This means that sometimes you need to run a command multiple times or distribute data-intensive commands over multiple cores. In this chapter, I have shown you how easy it is to parallelize commands. `parallel` is a very powerful and flexible tool to speed up ordinary command-line tools and distribute them. It offers a lot of functionality, and in this chapter I've only been able to scratch the surface. In the next chapter I'm going to cover the fourth step of the OSEMN model: modeling data.

For Further Exploration

Once you have a basic understanding of `parallel` and its most important options, I recommend that you take a look at [its online tutorial](#). Among other things, you'll learn different ways of specifying input, how to keep a log of all the jobs, and how to timeout, resume, and retry jobs. As Ole Tange, creator of `parallel`, says in this tutorial, “Your command line will love you for it.”

¹ Ole Tange, *parallel – Build and Execute Shell Command Lines from Standard Input in Parallel*, version 20161222, 2016, <https://www.gnu.org/software/parallel>.

² Eric B. Decker, James Youngman, and Kevin Dalley, *find – Search for Files in a Directory Hierarchy*, version 4.7.0, 2019, <https://www.gnu.org/software/findutils>.

³ Jeroen Janssens, *pbc – Parallel bc*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.

⁴ Amazon Web Services, *aws – Unified Tool to Manage AWS Services*, version 2.1.32, 2021, <https://aws.amazon.com/cli>.

- 5 Tatu Ylonen et al., *ssh – OpenSSH Remote Login Client*, version 1:8.2p1-4ubuntu0.2, 2020, <https://www.openssh.com>.
- 6 Peter Tobias, Bernd Eckenfels, and Michael Meskes, *hostname – Show or Set the System’s Host Name*, version 3.23, 2021, <https://sourceforge.net/projects/net-tools/>.
- 7 Antonio Diaz Diaz, *zcat – Decompress and Concatenate Files to Standard Output*, version 1.10, 2021, <https://www.nongnu.org/zutils/zutils.html>.

Chapter 9. Modeling Data

In this chapter we're going to perform the fourth step of the OSEMN model: modeling data. Generally speaking, a model is an abstract or higher-level description of your data. Modeling is a bit like creating visualizations in the sense that we're taking a step back from the individual data points to see the bigger picture.

Visualizations are characterized by shapes, positions, and colors: we can interpret them by looking at them. Models, on the other hand, are internally characterized by numbers, which means that computers can use them to do things like make predictions about new data points. (We can still visualize models so that we can try to understand them and see how they are performing.)

In this chapter I'll consider three types of algorithms commonly used to model data:

- Dimensionality reduction
- Regression
- Classification

These algorithms come from the field of statistics and machine learning, so I'm going to change the vocabulary a bit. Let's assume that I have a CSV file, also known as a *dataset*. Each row, except for the header, is considered a *data point*. Each data point has one or more *features*, or properties that have been measured. Sometimes a data point also has a *label*, which is, generally speaking, a judgment or outcome. This becomes more concrete when I introduce the wine dataset later in this chapter.

The first type of algorithm (dimensionality reduction) is most often unsupervised, which means that it creates a model based on the features of the dataset only. The other two types of algorithms (regression and

classification) are by definition supervised algorithms, which means that they also incorporate the labels into the model.

WARNING

This chapter is by no means an introduction to machine learning. That implies that I must skim over many details. My general advice is that you become familiar with an algorithm before applying it to your data. At the end of this chapter I recommend a few books about machine learning.

Overview

In this chapter, you'll learn how to:

- Reduce the dimensionality of your dataset using `tapkee`¹
- Predict the quality of white wine using `vw`²
- Classify wine as red or white using `skll`³

This chapter starts with the following file:

```
$ cd /data/ch09  
$ l  
total 4.0K  
-rw-r--r-- 1 dst dst 503 Jun 29 14:34 classify.cfg
```

The instructions for getting this file are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

More Wine, Please!

Throughout this chapter, I'll be using a dataset of wine tasters' notes on red and white varieties of a Portuguese wine called vinho verde. Each data point represents a wine. Each wine is rated on 11 physicochemical properties: (1) fixed acidity, (2) volatile acidity, (3) citric acid, (4) residual

sugar, (5) chlorides, (6) free sulfur dioxide, (7) total sulfur dioxide, (8) density, (9) pH, (10) sulfates, and (11) alcohol. There is also an overall quality score between 0 (very bad) and 10 (excellent), which is the median of at least three evaluations by wine experts. More information about this dataset is available at the [UCI Machine Learning Repository](#).

The dataset is split into two files: one for white wine and one for red wine. The first step is to obtain the two files using `curl` (and of course `parallel`, because I haven't got all day):

```
$ parallel "curl -sL http://archive.ics.uci.edu/ml/machine-learning-databases/wine-quality/winequality-{}.csv > wine-{}.csv" :::: red white
```

The triple colon is just another way to pass data to `parallel`:

```
$ cp /data/.cache/wine-*.csv _
```

Let's inspect both files and count the number of lines:

```
$ < wine-red.csv nl | ①
> fold | ②
> trim
    1 "fixed acidity";"volatile acidity";"citric acid";"residual
sugar";"chlor
ides";"free sulfur dioxide";"total sulfur
dioxide";"density";"pH";"sulphates";"a
lcohol";"quality"
    2 7.4;0.7;0;1.9;0.076;11;34;0.9978;3.51;0.56;9.4;5
    3 7.8;0.88;0;2.6;0.098;25;67;0.9968;3.2;0.68;9.8;5
    4 7.8;0.76;0.04;2.3;0.092;15;54;0.997;3.26;0.65;9.8;5
    5 11.2;0.28;0.56;1.9;0.075;17;60;0.998;3.16;0.58;9.8;6
    6 7.4;0.7;0;1.9;0.076;11;34;0.9978;3.51;0.56;9.4;5
    7 7.4;0.66;0;1.8;0.075;13;40;0.9978;3.51;0.56;9.4;5
    8 7.9;0.6;0.06;1.6;0.069;15;59;0.9964;3.3;0.46;9.4;5
... with 1592 more lines

$ < wine-white.csv nl | fold | trim
    1 "fixed acidity";"volatile acidity";"citric acid";"residual
sugar";"chlor
ides";"free sulfur dioxide";"total sulfur
dioxide";"density";"pH";"sulphates";"a
```

```

lcohol";"quality"
2 7;0.27;0.36;20.7;0.045;45;170;1.001;3;0.45;8.8;6
3 6.3;0.3;0.34;1.6;0.049;14;132;0.994;3.3;0.49;9.5;6
4 8.1;0.28;0.4;6.9;0.05;30;97;0.9951;3.26;0.44;10.1;6
5 7.2;0.23;0.32;8.5;0.058;47;186;0.9956;3.19;0.4;9.9;6
6 7.2;0.23;0.32;8.5;0.058;47;186;0.9956;3.19;0.4;9.9;6
7 8.1;0.28;0.4;6.9;0.05;30;97;0.9951;3.26;0.44;10.1;6
8 6.2;0.32;0.16;7;0.045;30;136;0.9949;3.18;0.47;9.6;6
... with 4891 more lines

$ wc -l wine-{red,white}.csv
1600 wine-red.csv
4899 wine-white.csv
6499 total

```

- ❶ For clarity, I use `nl` to add line numbers.
- ❷ To see the entire header, I use `fold`.

At first glance, this data appears to be quite clean. Still, let's scrub it so that it conforms more to what most command-line tools expect. Specifically, I will:

- Convert the header to lowercase
- Replace the semicolons with commas
- Replace spaces with underscores
- Remove unnecessary quotes

The tool `tr` can take care of all these things. Let's use a for loop—for old times' sake—to process both files:

```

$ for COLOR in red white; do
> < wine-$COLOR.csv tr '[A-Z]'; ' '[a-z],_` | tr -d \" > wine-${COLOR}-
clean.csv
> done

```

Let's also create a single dataset by combining the two files. I'll use `csvstack`⁴ to add a column named *type*, which will be "red" for rows of the first file and "white" for rows of the second file:

```
$ csvstack -g red,white -n type wine-{red,white}-clean.csv | ❶  
> xsv select 2-,1 > wine.csv ❷
```

- ❶ The new column *type* is placed at the beginning by `csvstack`.
- ❷ Some algorithms assume that the label is the last column, so I use `xsv` to move the column *type* to the end.

It's good practice to check whether there are any missing values in this dataset, because most machine learning algorithms can't handle them:

```
$ csvstat wine.csv --nulls  
1. fixed_acidity: False  
2. volatile_acidity: False  
3. citric_acid: False  
4. residual_sugar: False  
5. chlorides: False  
6. free_sulfur_dioxide: False  
7. total_sulfur_dioxide: False  
8. density: False  
9. ph: False  
10. sulphates: False  
11. alcohol: False  
12. quality: False  
13. type: False
```

Excellent! If there were any missing values, we could fill them with, say, the average or most common value of that feature. An alternative, less subtle approach is to remove any data points that have at least one missing value. Just out of curiosity, let's see what the distribution of quality looks like for both red and white wines (Figure 9-1):

```
$ rush run -t 'ggplot(df, aes(x = quality, fill = type)) + geom_density(adjust  
=
```

```
3, alpha = 0.5)' wine.csv > wine-quality.png
```

```
$ display wine-quality.png
```

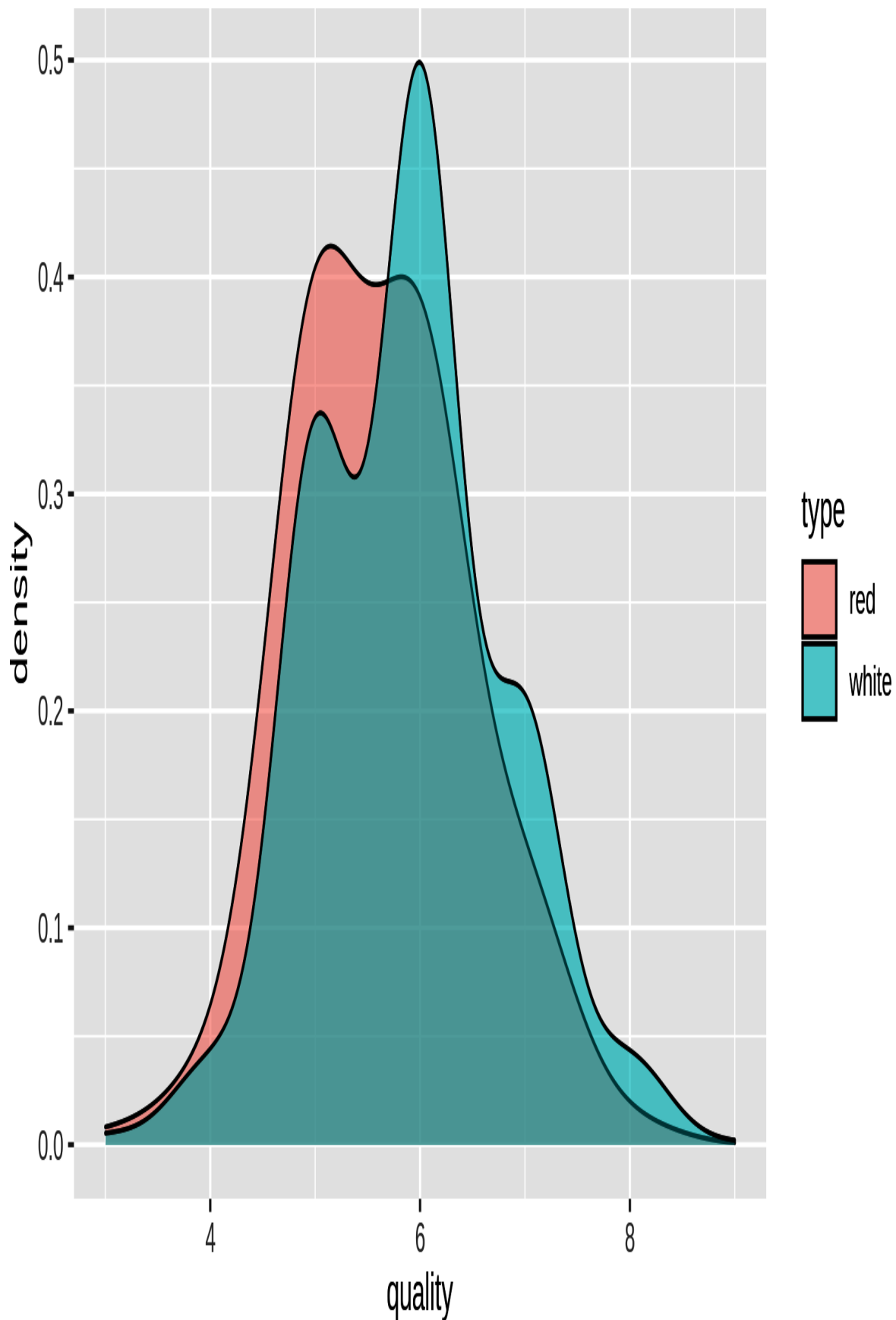


Figure 9-1. Comparing the quality of red and white wines using a density plot

From the density plot, you can see that the quality of white wine is distributed more toward higher values. Does this mean that white wines are better overall than red wines, or that the white wine experts more easily give higher scores than the red wine experts? That's something that the data doesn't tell us. Or is there perhaps a relationship between alcohol content and quality? Let's use `rush` to find out ([Figure 9-2](#)):

```
$ rush plot --x alcohol --y quality --color type --geom smooth wine.csv >
wine-a
lcohol-vs-quality.png

$ display wine-alcohol-vs-quality.png
```

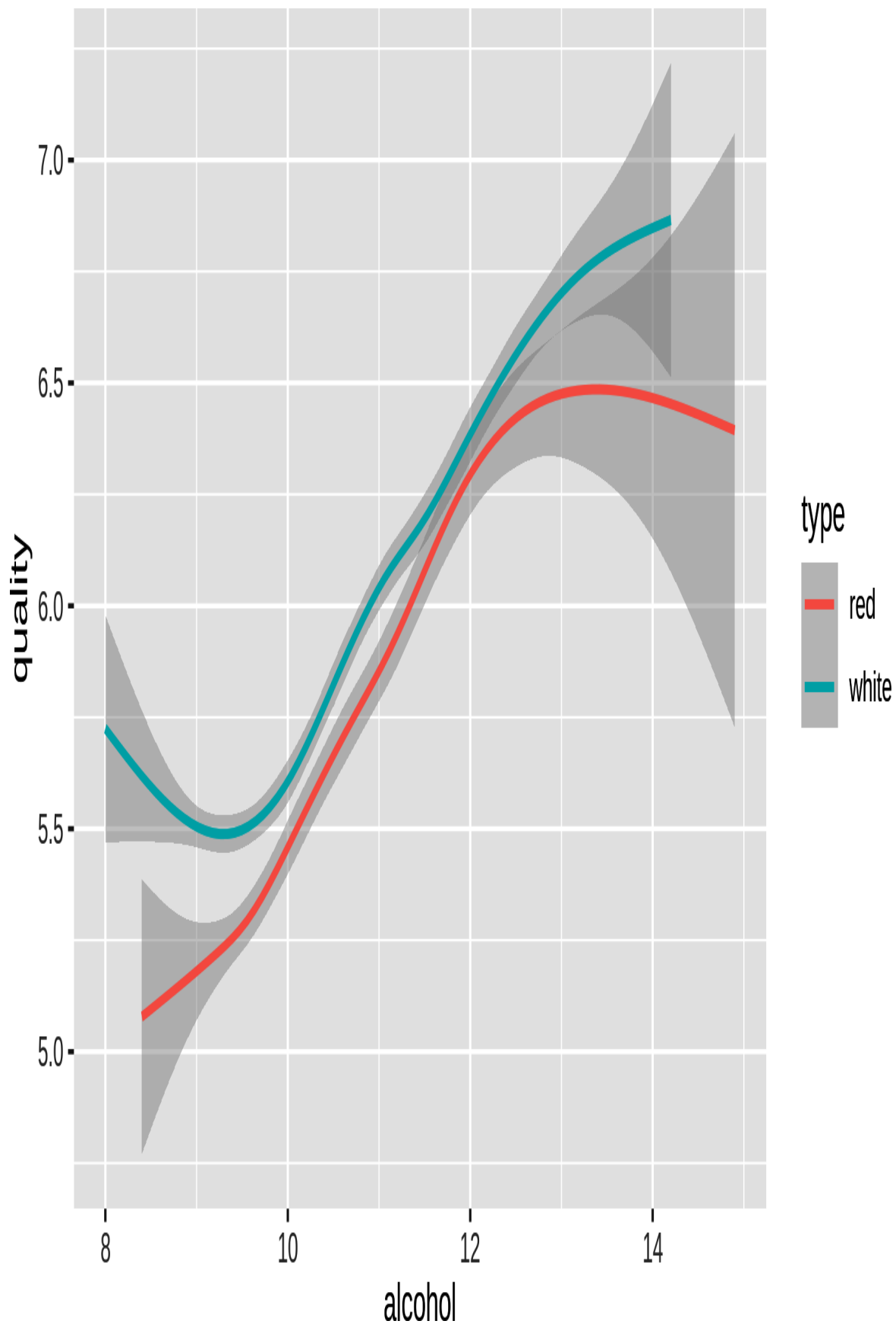


Figure 9-2. Relationship between the alcohol contents of wine and its quality

Eureka! Ahem, let's carry on with some modeling, shall we?

Dimensionality Reduction with Tapkee

The goal of dimensionality reduction is to map high-dimensional data points onto a lower-dimensional mapping. The challenge is to keep similar data points close together on the lower-dimensional mapping. As we've seen in the previous section, our wine dataset contains 13 features. I'll stick with two dimensions because that's straightforward to visualize.

Dimensionality reduction is often regarded as being part of exploration. It's useful for when there are too many features for plotting. You could do a scatter-plot matrix, but that shows you only two features at a time. It's also useful as a preprocessing step for other machine learning algorithms.

Most dimensionality reduction algorithms are unsupervised. This means they don't employ the labels of the data points to construct the lower-dimensional mapping.

In this section I'll look at two techniques used for dimensionality reduction: PCA, which stands for *Principal Component Analysis*,⁵ and t-SNE, which stands for *t-Distributed Stochastic Neighbor Embedding*.⁶

Introducing Tapkee

Tapkee is a C++ template library for dimensionality reduction.⁷ The library contains implementations of many dimensionality-reduction algorithms, including:

- Locally Linear Embedding
- Isomap
- Multidimensional Scaling
- PCA

- t-SNE

More information about these algorithms can be found on [Tapkee's website](#). Although Tapkee is mainly a library that can be included in other applications, it also offers the command-line tool `tapkee`. I'll use this tool to perform dimensionality reduction on our wine dataset.

Linear and Nonlinear Mappings

First, I'll scale the features using standardization so that each feature is equally important. This generally leads to better results when applying machine learning algorithms.

To scale, I use `rush` and the tidyverse package:

```
$ rush run --tidyverse --output wine-scaled.csv \
> 'select(df, -type) %>% ❶
> scale() %>% ❷
> as_tibble() %>% ❸
> mutate(type = df$type)' wine.csv ❹

$ csvlook wine-scaled.csv
fixed_acidity | volatile_acidity | citric_acid | residual_sugar | chlorides
...
...
...
0.142... | 2.189... | -2.193... | -0.745... | ...
0.451... | 3.282... | -2.193... | -0.598... | ...
0.451... | 2.553... | -1.917... | -0.661... | ...
3.074... | -0.362... | 1.661... | -0.745... | ...
0.142... | 2.189... | -2.193... | -0.745... | ...
0.142... | 1.946... | -2.193... | -0.766... | ...
0.528... | 1.581... | -1.780... | -0.808... | ...
0.065... | 1.885... | -2.193... | -0.892... | ...
... with 6489 more lines
```

- ❶ I need to temporarily remove the column `type` because `scale()` only works on numerical columns.
- ❷ The `scale()` function accepts a data frame but returns a matrix.

- ③ The function `as_tibble()` converts the matrix back to a data frame.
- ④ Finally, I add back the *type* column.

Now we apply both dimensionality-reduction techniques and visualize the mapping using `Rio-scatter`:

```
$ xsv select '!type' wine-scaled.csv | ①
> header -d | ②
> tapkee --method pca | ③
> tee wine-pca.txt | trim
-0.568882,3.34818
-1.19724,3.22835
-0.952507,3.23722
-1.60046,1.67243
-0.568882,3.34818
-0.556231,3.15199
-0.53894,2.28288
1.104,2.56479
0.231315,2.86763
-1.18363,1.81641
... with 6487 more lines
```

- ① Deselect the column *type*.
- ② Remove the header.
- ③ Apply PCA:

```
$ < wine-pca.txt header -a pc1,pc2 | ①
> paste -d, - <(xsv select type wine-scaled.csv) | ②
> tee wine-pca.csv | csvlook
| pc1 | pc2 | type |
| -0.569... | 3.348... | red |
| -1.197... | 3.228... | red |
| -0.953... | 3.237... | red |
| -1.600... | 1.672... | red |
```

```
| -0.569... | 3.348... | red |
| -0.556... | 3.152... | red |
| -0.539... | 2.283... | red |
| 1.104...  | 2.565... | red |
... with 6489 more lines
```

- ① Add back the header with columns $pc1$ and $pc2$.
- ② Add back the column $type$.

Now we can create a scatter plot (Figure 9-3):

```
$ rush plot --x pc1 --y pc2 --color type --shape type wine-pca.csv > wine-
pca.png
g

$ display wine-pca.png
```

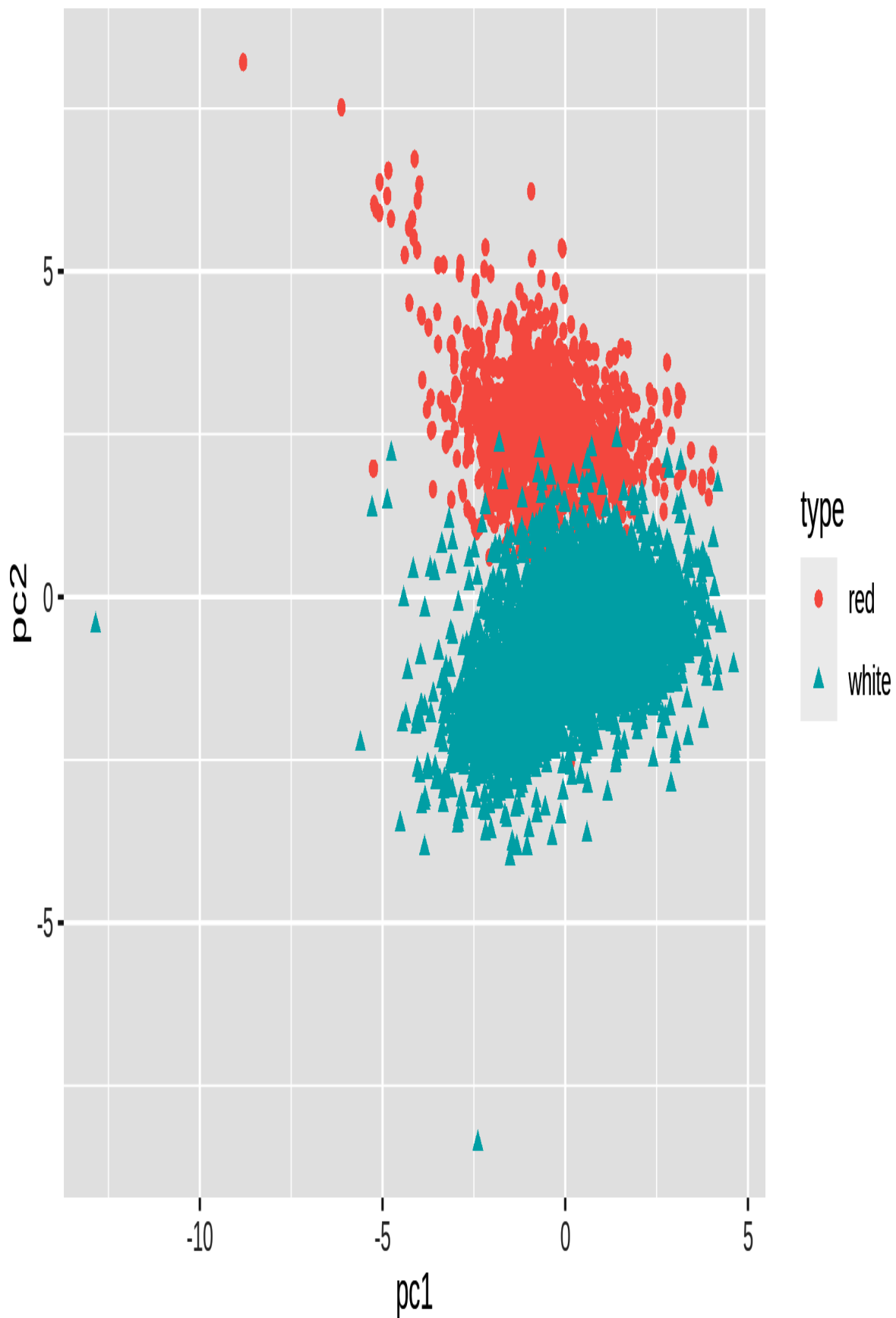


Figure 9-3. Linear dimensionality reduction with PCA

Let's perform t-SNE with the same approach:

```
$ xsv select '!type' wine-scaled.csv | ❶
> header -d | ❷
> tapkee --method t-sne | ❸
> header -a x,y | ❹
> paste -d, - <(xsv select type wine-scaled.csv) | ❺
> rush plot --x x --y y --color type --shape type > wine-tsne.png ❻
```

- ❶ Deselect the column *type*.
- ❷ Remove the header.
- ❸ Apply t-SNE.
- ❹ Add back the header with columns *x* and *y*.
- ❺ Add back the column *type*.
- ❻ Create a scatter plot with the same approach (Figure 9-4):

```
$ display wine-tsne.png
```

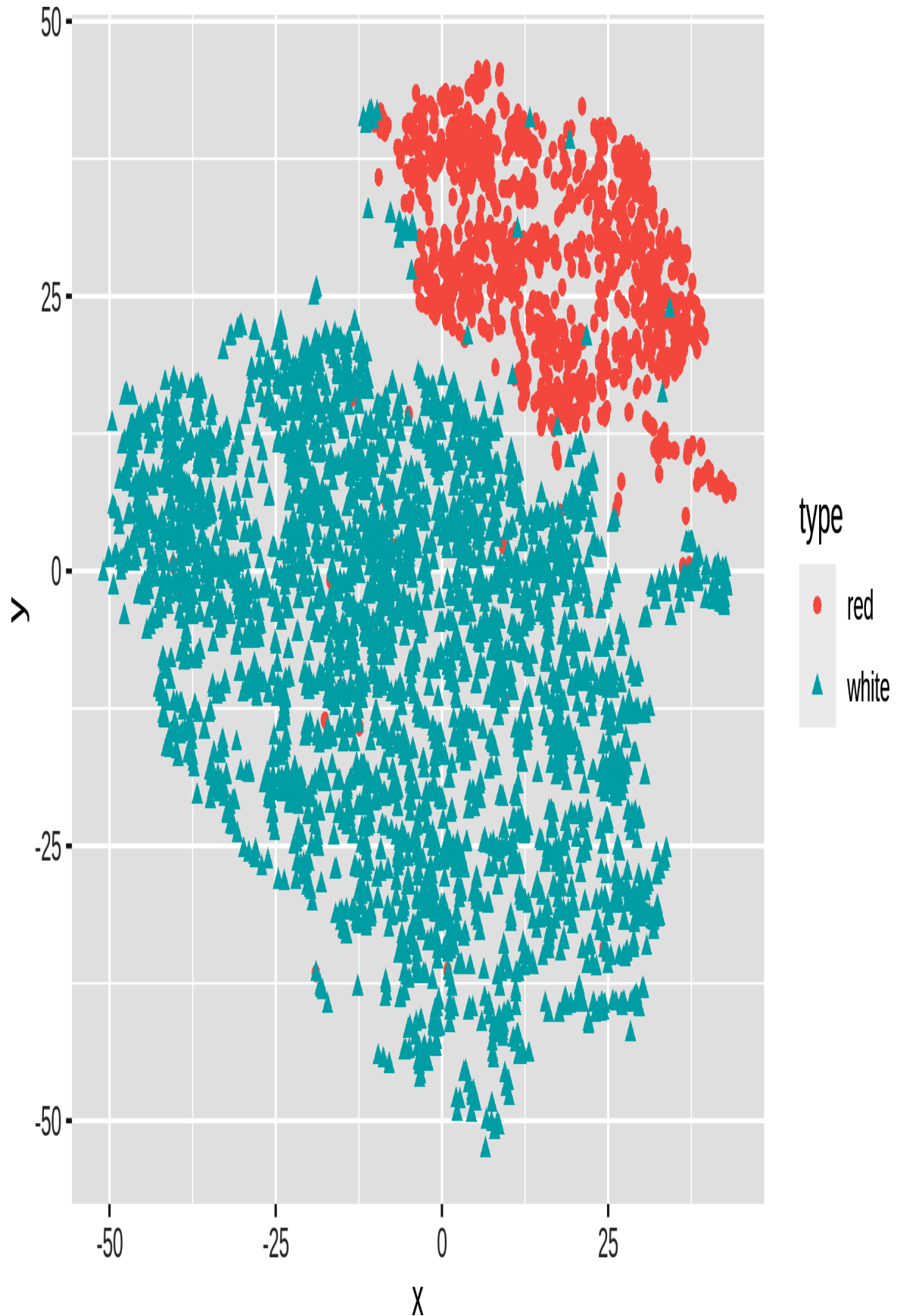


Figure 9-4. Nonlinear dimensionality reduction with t-SNE

We can see that t-SNE does a better job than PCA at separating the red and white wines based on their physicochemical properties. These scatter plots verify that the dataset has a certain structure; there's a relationship between the features and the labels. Knowing this, I'm comfortable moving forward by applying supervised machine learning. I'll start with a regression task and then continue with a classification task.

Regression with Vowpal Wabbit

In this section, I'm going to create a model that predicts the quality of the white wine, based on its physicochemical properties. Because the quality is a number between 0 and 10, we can consider this a regression task.

For this I'll be using Vowpal Wabbit, or `vw`.

Preparing the Data

Instead of working with CSV, `vw` has its own data format. The tool `csv2vw`⁸ can, as its name implies, convert CSV to this format. The `--label` option is used to indicate which column contains the labels. Let's examine the result:

```
$ csv2vw wine-white-clean.csv --label quality | trim
6 | alcohol:8.8 chlorides:0.045 citric_acid:0.36 density:1.001 fixed_acidity:7
...
6 | alcohol:9.5 chlorides:0.049 citric_acid:0.34 density:0.994
fixed_acidity:6...
6 | alcohol:10.1 chlorides:0.05 citric_acid:0.4 density:0.9951
fixed_acidity:8...
6 | alcohol:9.9 chlorides:0.058 citric_acid:0.32 density:0.9956
fixed_acidity:7...
6 | alcohol:9.9 chlorides:0.058 citric_acid:0.32 density:0.9956
fixed_acidity:7...
6 | alcohol:10.1 chlorides:0.05 citric_acid:0.4 density:0.9951
fixed_acidity:8...
6 | alcohol:9.6 chlorides:0.045 citric_acid:0.16 density:0.9949
fixed_acidity:6...
6 | alcohol:8.8 chlorides:0.045 citric_acid:0.36 density:1.001 fixed_acidity:7
...
```

```
6 | alcohol:9.5 chlorides:0.049 citric_acid:0.34 density:0.994  
fixed_acidity:6....  
6 | alcohol:11 chlorides:0.044 citric_acid:0.43 density:0.9938  
fixed_acidity:8....  
... with 4888 more lines
```

In this format, each line is one data point. The line starts with the label, followed by a pipe symbol and then by feature name/value pairs separated by spaces. While this format may seem overly verbose when compared to the CSV format, it does offer more flexibility, such as weights, tags, namespaces, and a sparse feature representation. With the wine dataset, we don't need this flexibility, but it might be useful when applying `vw` to more complicated problems. [This article](#) explains the `vw` format in more detail.

Once we've created, or *trained*, a regression model, it can be used to make predictions about new, unseen data points. In other words, if we give the model a wine it hasn't seen before, it can predict, or *test*, its quality. To properly evaluate the accuracy of these predictions, we need to set aside some data that will not be used for training. It's common to use 80% of the complete dataset for training and the remaining 20% for testing.

I can do this by first splitting the complete dataset into five equal parts using `split`.⁹ I verify the number of data points in each part using `wc`:

```
$ csv2vw wine-white-clean.csv --label quality |  
> shuf | ①  
> split -d -n r/5 - wine-part-  
  
$ wc -l wine-part-*  
980 wine-part-00  
980 wine-part-01  
980 wine-part-02  
979 wine-part-03  
979 wine-part-04  
4898 total
```

- ① The tool `shuf`¹⁰ randomizes the dataset to ensure that both the training and the test have similar quality distribution.

Now I can use the first part (so 20%) for the testing set *wine-test.vw* and combine the four remaining parts (i.e., 80%) into the training set *wine-train.vw*:

```
$ mv wine-part-00 wine-test.vw  
$ cat wine-part-* > wine-train.vw  
$ rm wine-part-*  
$ wc -l wine-*.vw  
    980 wine-test.vw  
  3918 wine-train.vw  
  4898 total
```

Now we're ready to train a model using *vw*.

Training the Model

The tool *vw* accepts many different options (nearly four hundred!). Luckily, you don't need all of them to be effective. To annotate the options I use here, I'll put each one on a separate line:

```
$ vw \  
> --data wine-train.vw \ ①  
> --final_regressor wine.model \ ②  
> --passes 10 \ ③  
> --cache_file wine.cache \ ④  
> --nn 3 \ ⑤  
> --quadratic :: \ ⑥  
> --l2 0.000005 \ ⑦  
> --bit_precision 25 ⑧  
creating quadratic features for pairs: ::  
WARNING: any duplicate namespace interactions will be removed  
You can use --leave_duplicate_interactions to disable this behaviour.  
using l2 regularization = 5e-06  
final_regressor = wine.model  
Num weight bits = 25  
learning rate = 0.5  
initial_t = 0  
power_t = 0.5  
decay_learning_rate = 1  
creating cache_file = wine.cache
```

```

Reading datafile = wine-train.vw
num sources = 1
Enabled reductions: gd, generate_interactions, nn, scorer
average since      example      example current current current
loss   last       counter      weight    label predict features
49.000000 49.000000          1           1.0    7.0000  0.0000    78
38.224279 27.448559          2           2.0    6.0000  0.7609    78
29.787616 21.350952          4           4.0    6.0000  1.5382    78
23.172403 16.557191          8           8.0    6.0000  2.2755    78
16.745619 10.318835         16          16.0   5.0000  3.0704    78
11.383260 6.020901          32          32.0   6.0000  3.9637    78
6.953757 2.524254          64          64.0   5.0000  4.9483    78
4.096179 1.238601          128         128.0  7.0000  5.6035    78
2.365188 0.634197          256         256.0  5.0000  6.0535    78
1.575675 0.786162          512         512.0  5.0000  6.0247    78
1.210981 0.846286          1024        1024.0 5.0000  5.7023    78
0.977191 0.743401          2048        2048.0 7.0000  6.0133    78
0.936266 0.936266          4096        4096.0 6.0000  5.7761    78 h
0.819179 0.702092          8192        8192.0 6.0000  5.9635    78 h
0.722494 0.625809          16384       16384.0 7.0000  6.2734    78 h
0.668416 0.614337          32768       32768.0 6.0000  5.0941    78 h

finished run
number of examples per pass = 3527
passes used = 10
weighted example sum = 35270.000000
weighted label sum = 206890.000000
average loss = 0.605274 h
best constant = 5.865891
total feature number = 2749740

```

- ❶ The file *wine-train.vw* is used to train the model.
- ❷ The model, or *regressor*, will be stored in the file *wine.model*.
- ❸ The learning algorithm will make 10 passes over the training data.
- ❹ Caching is needed when making multiple passes.
- ❺ Use a neural network with three hidden units.

- ⑥ Create and use quadratic features, based on all input features. Any duplicates will be removed by `vw`.
- ⑦ Use l2 regularization.
- ⑧ Use 25 bits to store the features.

Now that I have trained a regression model, let's use it to make predictions.

Testing the Model

The model is stored in the file `wine.model`. To use that model to make predictions, I run `vw` again, but now with a different set of options:

```
$ vw \
> --data wine-test.vw \ ①
> --initial_regressor wine.model \ ②
> --testonly \ ③
> --predictions predictions \ ④
> --quiet ⑤

$ bat predictions | trim
5.963508
6.166912
5.336860
6.251552
6.034288
6.529794
5.025755
6.123046
6.090032
5.573547
... with 970 more lines
```

- ① The file `wine-test.vw` is used to test the model.
- ② Use the model stored in the file `wine.model`.

- ③ Ignore label information and just test.
- ④ The predictions are stored in a file called *predictions*.
- ⑤ Don't output diagnostics or progress updates.

Let's use `paste` to combine the predictions in the file *predictions* with the true, or *observed*, values that are in the file *wine-test.vw*. Using `awk`, I can compare the predicted values with the observed values and compute the mean absolute error (MAE). The MAE tells us how far off `vw` is on average when it comes to predicting the quality of a white wine:

```
$ paste -d, predictions <(cut -d '|' -f 1 wine-test.vw) |  
> tee results.csv |  
> awk -F, '{E+=sqrt((\$1-\$2)^2)} END {print "MAE: " E/NR}' |  
> cowsay ❶  
  
-----  
< MAE: 0.569113 >  
-----  
 \ ^__^  
  \  (oo)\_____  
   (__)\       )\/\|  
    ||----w |  
    ||     ||
```

So the predictions are, on average, about 0.6 points off. Let's visualize the relationship between the observed values and the predicted values using `rush plot` (Figure 9-5):

```
$ < results.csv header -a "predicted,observed" |  
> rush plot --x observed --y predicted --geom jitter > wine-regression.png  
  
$ display wine-regression.png
```

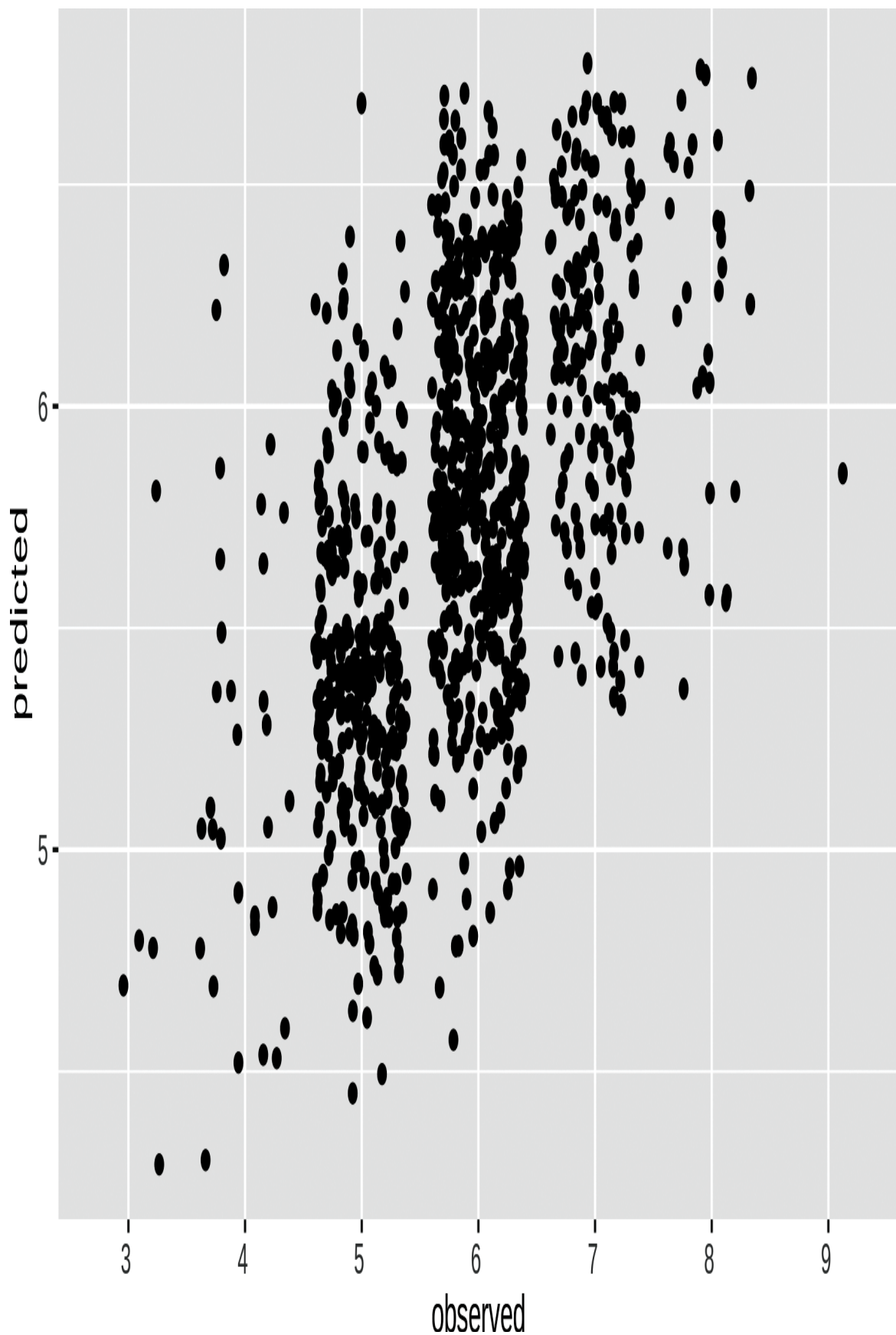


Figure 9-5. Regression with Vowpal Wabbit

I can imagine that the options used to train the model might be a bit overwhelming. Let's see how `vw` performs when I use all the default values:

```
$ vw -d wine-train.vw -f wine2.model --quiet ①  
$ vw -data wine-test.vw -i wine2.model -t -p predictions --quiet ②  
$ paste -d, predictions <(cut -d ' '| -f 1 wine-test.vw) | ③  
> awk -F, '{E+=sqrt((\$1-\$2)^2)} END {print "MAE: " E/NR}'  
MAE: 0.626015
```

- ① Train a regression model.
- ② Test the regression model.
- ③ Compute the MAE.

Apparently, with the default values the MAE is 0.046 higher, meaning that the predictions are slightly worse.

In this section, I've only been able to scratch the surface of what `vw` can do. There's a reason why it accepts so many options. Besides regression, it also supports binary classification, multiclass classification, reinforcement learning, and Latent Dirichlet Allocation, among other things. You can learn more from the many tutorials and articles on the [Vowpal Wabbit website](#).

Classification with SciKit-Learn Laboratory

In this section I'm going to train a classification model, or *classifier*, that predicts whether a wine is red or white. While we could use `vw` for this, I'd like to demonstrate another tool: SciKit-Learn Laboratory (SKLL). As the name implies, it's built on top of SciKit-Learn, a popular machine learning package for Python. SKLL, itself a Python package, provides the

`run_experiment` tool, which makes it possible to use SciKit-Learn from the command line. Instead of `run_experiment`, I use the alias `skll` because I find it easier to remember, as it corresponds to the package name:

```
$ alias skll=run_experiment  
  
$ skll  
usage: run_experiment [-h] [-a NUM_FEATURES] [-A] [-k] [-l] [-m MACHINES]  
                      [-q QUEUE] [-r] [-v] [--version]  
                      config_file [config_file ...]  
run_experiment: error: the following arguments are required: config_file
```

Preparing the Data

`skll` expects the training and test datasets to have the same filenames, located in separate directories. Because its predictions are not necessarily in the same order as the original dataset, I add a column, *id*, that contains a unique identifier so that I can match the predictions with the correct data points. Let's create a balanced dataset:

```
$ NUM_RED="$(wine-red-clean.csv wc -l)" ❶  
  
$ csvstack -n type -g red,white \\ ❷  
  > wine-red-clean.csv \\  
  > <(< wine-white-clean.csv body shuf | head -n $NUM_RED) |  
  > body shuf |  
  > nl -s, -w1 -v0 | ❸  
  > sed '1s/0,/id,' | ❹  
  > tee wine-balanced.csv | csvlook  
|   id | type | fixed_acidity | volatile_acidity | citric_acid |  
residual_sug...  
|---|---|---|---|---|  
| ... | | | | |  
| 1 | 1 | red | 7.50 | 0.400 | 0.18 |  
| 1... | | | | |  
| 2 | 2 | red | 6.70 | 0.760 | 0.02 |  
| 1... | | | | |  
| 3 | 3 | white | 5.70 | 0.260 | 0.27 |  
| 4... | | | | |  
| 4 | 4 | white | 5.80 | 0.230 | 0.27 |  
| 1... | | | | |  
| 5 | 5 | red | 7.90 | 0.765 | 0.00 |  
| 2... | | | | |
```

```

|      6 | white |      5.30 |      0.260 |      0.23 |
5...
|      7 | red   |      6.70 |      0.580 |      0.08 |
1...
|      8 | white |      8.00 |      0.170 |      0.29 |
2...
... with 3190 more lines

```

- ❶ Store the number of red wines in the variable NUM_RED.
- ❷ Combine all red wines with a random sample of white wines.
- ❸ Add line numbers using `nl` in front of each line.
- ❹ Replace the `0` on the first line with `id` so that it's a proper column name.

Let's split this balanced dataset into a training set and a test set:

```

$ mkdir -p {train,test}

$ HEADER=$(< wine-balanced.csv header)

$ < wine-balanced.csv header -d | shuf | split -d -n r/5 - wine-part-

$ wc -l wine-part-*
 640 wine-part-00
 640 wine-part-01
 640 wine-part-02
 639 wine-part-03
 639 wine-part-04
 3198 total

$ cat wine-part-00 | header -a $HEADER > test/features.csv && rm wine-part-00

$ cat wine-part-* | header -a $HEADER > train/features.csv && rm wine-part-*

$ wc -l t*/features.csv
 641 test/features.csv
 2559 train/features.csv
 3200 total

```

Now that I have a balanced training dataset and a balanced test dataset, I can continue with building a classifier.

Running the Experiment

Training a classifier in `skll` is done by defining an experiment in a configuration file. It consists of several sections that specify, for example, where to look for the datasets, which classifiers to train, and what kind of output to generate. Here's the configuration file `classify.cfg` that I'll use:

```
$ bat classify.cfg
-----
| File: classify.cfg
-----
|
1 | [General]
2 | experiment_name = wine
3 | task = evaluate
4 |
5 | [Input]
6 | train_directory = train
7 | test_directory = test
8 | featuresets = [["features"]]
9 | feature_scaling = both
10 | label_col = type
11 | id_col = id
12 | shuffle = true
13 | learners = ["KNeighborsClassifier", "LogisticRegression",
"DecisionTree
    | Classifier", "RandomForestClassifier"]
14 | suffix = .csv
15 |
16 | [Tuning]
17 | grid_search = false
18 | objectives = ["neg_mean_squared_error"]
19 | param_grids = [ {}, {}, {}, {} ]
20 |
21 | [Output]
22 | logs = output
23 | results = output
24 | predictions = output
25 | models = output
```

I run the experiment using `skll`:

```
$ skll -l classify.cfg 2>/dev/null
```

The option `-l` specifies to run in local mode. `skll` also offers the possibility to run experiments on clusters. The time it takes to run an experiment depends on the complexity of the chosen algorithms and the size of the data.

Parsing the Results

Once all classifiers have been trained and tested, the results can be found in the directory `output`:

```
$ ls -1 output
wine_features_DecisionTreeClassifier.log
wine_features_DecisionTreeClassifier.model
wine_features_DecisionTreeClassifier_predictions.tsv
wine_features_DecisionTreeClassifier.results
wine_features_DecisionTreeClassifier.results.json
wine_features_KNeighborsClassifier.log
wine_features_KNeighborsClassifier.model
wine_features_KNeighborsClassifier_predictions.tsv
wine_features_KNeighborsClassifier.results
wine_features_KNeighborsClassifier.results.json
wine_features_LogisticRegression.log
wine_features_LogisticRegression.model
wine_features_LogisticRegression_predictions.tsv
wine_features_LogisticRegression.results
wine_features_LogisticRegression.results.json
wine_features_RandomForestClassifier.log
wine_features_RandomForestClassifier.model
wine_features_RandomForestClassifier_predictions.tsv
wine_features_RandomForestClassifier.results
wine_features_RandomForestClassifier.results.json
wine.log
wine_summary.tsv
```

`skll` generates four files for each classifier: one log, two files with results, and one file with predictions. I extract the algorithm names and sort them by their accuracies using the following SQL query:

```
$ < output/wine_summary.tsv csvsql --query "SELECT learner_name, accuracy FROM
  s
  tdn ORDER BY accuracy DESC" | csvlook -I
  | learner_name      | accuracy   |
  |-----+-----|
  | RandomForestClassifier | 0.9890625 |
  | KNeighborsClassifier    | 0.9875     |
  | LogisticRegression     | 0.9859375 |
  | DecisionTreeClassifier  | 0.9640625 |
```

The relevant column here is *accuracy*, which indicates the percentage of data points that are classified correctly. From this we see that actually all algorithms are performing really well. `RandomForestClassifier` comes out as the best-performing algorithm, closely followed by `KNeighborsClassifier`.

Each JSON file contains a confusion matrix, giving you additional insight into the performance of each classifier. A confusion matrix is a table in which the columns refer to the true labels (red and white) and the rows refer to the predicted labels. Higher numbers on the diagonal mean more correct predictions. With `jq` I can print the name of each classifier and extract the associated confusion matrix:

```
$ jq -r '.[] | "\(.learner_name):\n\(.result_table)\n"' output/*.json
DecisionTreeClassifier:
+-----+-----+-----+-----+-----+
|     | red | white | Precision | Recall | F-measure |
+=====+=====+=====+=====+=====+
| red | [309] |     8 |      0.954 |    0.975 |     0.964 |
+-----+-----+-----+-----+-----+
| white |    15 | [308] |      0.975 |    0.954 |     0.964 |
+-----+-----+-----+-----+-----+
(row = reference; column = predicted)

KNeighborsClassifier:
+-----+-----+-----+-----+-----+
|     | red | white | Precision | Recall | F-measure |
```

```

+=====+=====+=====+=====+=====+
|   red | [311] |     6 |    0.994 |    0.981 |    0.987 |
+-----+-----+-----+-----+-----+
| white |     2 | [321] |    0.982 |    0.994 |    0.988 |
+-----+-----+-----+-----+

```

(row = reference; column = predicted)

LogisticRegression:

```

+-----+-----+-----+-----+
|       |   red |   white | Precision | Recall | F-measure |
+=====+=====+=====+=====+=====+
|   red | [311] |     6 |    0.990 |    0.981 |    0.986 |
+-----+-----+-----+-----+
| white |     3 | [320] |    0.982 |    0.991 |    0.986 |
+-----+-----+-----+-----+

```

(row = reference; column = predicted)

RandomForestClassifier:

```

+-----+-----+-----+-----+
|       |   red |   white | Precision | Recall | F-measure |
+=====+=====+=====+=====+=====+
|   red | [313] |     4 |    0.991 |    0.987 |    0.989 |
+-----+-----+-----+-----+
| white |     3 | [320] |    0.988 |    0.991 |    0.989 |
+-----+-----+-----+-----+

```

(row = reference; column = predicted)

A confusion matrix is especially helpful when you have more than two classes, so that you can see which kinds of misclassifications happen, and when the cost of an incorrect classification is not the same for each class.

From a usage perspective, it's interesting to consider that `vw` and `skll` take two different approaches: `vw` uses command-line options, whereas `skll` requires a separate file. Both approaches have their advantages and disadvantages. While command-line options enable more ad hoc usage, a configuration file is perhaps easier to reproduce. Then again, as we've seen, invoking `vw` with any number of options can easily be placed in script or in a *Makefile*. The opposite approach—making `skll` accept options such that it doesn't need a configuration file—is less straightforward.

Summary

In this chapter we've looked at modeling data. Through examples, I dived into three different machine learning tasks—namely dimensionality reduction, which is unsupervised, and regression and classification, which are both supervised. A proper machine learning tutorial is unfortunately beyond the scope of this book. In the next section I have a couple of recommendations in case you want to learn more about machine learning. This was the fourth and last step of the OSEMN model for data science that I'm covering in this book. The next chapter is the last intermezzo chapter and will be about leveraging the command line elsewhere.

For Further Exploration

- The book *Python Machine Learning* by Sebastian Raschka and Vahid Mirjalili (Packt) offers a comprehensive overview of machine learning and how to apply it using Python.
- The later chapters of *R for Everyone* by Jared Lander (Addison-Wesley) explain how to accomplish various machine learning tasks using R.
- If you want to get a deeper understanding of machine learning, I highly recommend you pick up *Pattern Recognition and Machine Learning* by Christopher Bishop (Springer) and *Information Theory, Inference, and Learning Algorithms* by David MacKay (Cambridge University Press).
- If you're interested in learning more about the t-SNE algorithm, I recommend the original article about it: “[Visualizing Data Using t-SNE](#)” by Laurens van der Maaten and Geoffrey Hinton.

¹ Sergey Lisitsyn, Christian Widmer, and Fernando J. Iglesias Garcia, *tapkee – an Efficient Dimension Reduction Library*, version 1.2, 2013, <http://tapkee.lisitsyn.me>.

- 2 John Langford, *vw – Fast Machine Learning Library for Online Learning*, version 8.10.1, 2021, <https://vowpalwabbit.org>.
- 3 Educational Testing Service, *skll – SciKit-Learn Laboratory*, version 2.5.0, 2021, <https://skll.readthedocs.org>.
- 4 Christopher Groskopf, *csvstack – Stack Up the Rows from Multiple CSV Files*, version 1.0.5, 2020, <https://csvkit.rtfd.org>.
- 5 Karl Pearson, “On Lines and Planes of Closest Fit to Systems of Points in Space,” *Philosophical Magazine* 2, no. 11 (1901): 559–72, <http://pca.narod.ru/pearson1901.pdf>.
- 6 Laurens van der Maaten and Geoffrey Everest Hinton, “Visualizing Data Using t-SNE,” *Journal of Machine Learning Research* 9 (2008): 2579–605, <https://www.jmlr.org/papers/volume9/vandermaaten08a/vandermaaten08a.pdf>.
- 7 Sergey Lisitsyn, Christian Widmer, and Fernando J. Iglesias Garcia, “Tapkee: An Efficient Dimension Reduction Library,” *Journal of Machine Learning Research* 14 (2013): 2355–59, <https://jmlr.org/papers/volume14/lisitsyn13a/lisitsyn13a.pdf>.
- 8 Jeroen Janssens, *csv2vw – Convert CSV to Vowpal Wabbit Format*, version 0.1, 2021, <https://github.com/jeroenjanssens/dsutils>.
- 9 Torbjorn Granlund and Richard M. Stallman, *split – Split a File into Pieces*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.
- 10 Paul Eggert, *shuf – Generate Random Permutations*, version 8.30, 2019, <https://www.gnu.org/software/coreutils>.

Chapter 10. Polyglot Data Science

A *Polyglot* is someone who speaks multiple languages. A polyglot data scientist, as I see it, is someone who uses multiple programming languages, tools, and techniques to obtain, scrub, explore, and model data.

The command line stimulates a polyglot approach. The command line doesn't care which programming language a tool is written in, as long as it adheres to the Unix philosophy. We saw that very clearly in [Chapter 4](#), where we created command-line tools in Bash, Python, and R. Moreover, we executed SQL queries directly on CSV files and executed R expressions from the command line. In short, we have already been doing polyglot data science without fully realizing it!

In this chapter I'm going take this further by flipping it around. I'm going to show you how to leverage the command line from various programming languages and environments. Because let's be honest: we're not going to spend our entire data science careers at the command line. As for me, when I'm analyzing some data, I often use the RStudio integrated development environment (IDE); and when I'm implementing something, I often use Python. I use whatever helps me get the job done.

I find it comforting to know that the command line is often within arm's reach, without my having to switch to a different application. It allows me to quickly run a command without switching to a separate application and breaking my workflow. Examples are downloading files with `curl`, inspecting a piece of data with `head`, creating a backup with `git`, and compiling a website with `make`. Generally speaking, these are tasks that normally require a lot of code or simply cannot be done at all without the command line.

Overview

In this chapter, you'll learn how to:

- Run a terminal within JupyterLab and RStudio IDE
- Interact with arbitrary command-line tools in Python and R
- Transform data using shell commands in Apache Spark

This chapter starts with the following files:

```
$ cd /data/ch10  
$ l  
total 176K  
-rw-r--r-- 1 dst dst 164K Jun 29 14:36 alice.txt  
-rwxr-xr-x 1 dst dst  408 Jun 29 14:36 count.py*  
-rw-r--r-- 1 dst dst  460 Jun 29 14:36 count.R  
-rw-r--r-- 1 dst dst 1.7K Jun 29 14:36 Untitled1337.ipynb
```

The instructions for getting these files are in [Chapter 2](#). Any other files are either downloaded or generated using command-line tools.

Jupyter

Project Jupyter is an open source project, born out of the IPython project in 2014 as it evolved to support interactive data science and scientific computing across all programming languages. Jupyter supports more than 40 programming languages, including Python, R, Julia, and Scala. In this section I'll focus on Python.

The project includes JupyterLab, Jupyter Notebook, and Jupyter Console. I'll start with Jupyter Console, as it is the most basic option for working with Python in an interactive way. Here's a Jupyter Console session illustrating a couple of ways to leverage the command line:

```
$ jupyter console  
Jupyter console 6.4.0
```

```
Python 3.9.4 (default, Apr  4 2021, 19:38:44)
Type 'copyright', 'credits' or 'license' for more information
IPython 7.23.0 -- An enhanced Interactive Python. Type '?' for help.
```

```
In [1]: ! date ❶
Sun May  2 01:45:06 PM CEST 2021
```

```
In [2]: ! pip install --upgrade requests
Requirement already satisfied: requests in
/home/dst/.local/lib/python3.9/site-p
ackages (2.25.1)
Collecting requests
  Using cached requests-2.25.1-py2.py3-none-any.whl (61 kB)
  Downloading requests-2.25.0-py2.py3-none-any.whl (61 kB)
    |██████████| 61 kB 2.1 MB/s
Requirement already satisfied: urllib3<1.27,>=1.21.1 in
/home/dst/.local/lib/pyt
hon3.9/site-packages (from requests) (1.26.4)
Requirement already satisfied: certifi>=2017.4.17 in
/home/dst/.local/lib/python
3.9/site-packages (from requests) (2020.12.5)
Requirement already satisfied: chardet<5,>=3.0.2 in /usr/lib/python3/dist-
packag
es (from requests) (4.0.0)
Requirement already satisfied: idna<3,>=2.5 in
/home/dst/.local/lib/python3.9/si
te-packages (from requests) (2.10)
```

```
In [3]: ! head alice.txt
Project Gutenberg's Alice's Adventures in Wonderland, by Lewis Carroll
```

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Title: Alice's Adventures in Wonderland

```
In [4]: len(open("alice.txt").read().strip().split("\n")) ❷
Out[4]: 3735
```

```
In [5]: total_lines = ! < alice.txt wc -l
```

```
In [6]: total_lines
Out[6]: ['3735']
```

```
In [7]: int(total_lines[0]) ③
Out[7]: 3735

In [8]: url = "https://www.gutenberg.org/files/11/old/11.txt"

In [9]: import requests ④

In [10]: with open("alice2.txt", "wb") as f:
....:     response = requests.get(url)
....:     f.write(response.content)
....:

In [11]: ! curl '{url}' > alice3.txt ⑤
% Total    % Received % Xferd  Average Speed   Time     Time     Time
Current                                         Dload  Upload   Total   Spent   Left  Speed
100  163k  100  163k    0      0  211k      0 --::-- --::-- --::-- 211k

In [12]: ! ls alice*.txt
alice2.txt  alice3.txt  alice.txt

In [13]: ! rm -v alice{2,3}.txt ⑥
zsh:1: no matches found: alice(2, 3).txt

In [14]: ! rm -v alice{{2,3}}.txt
removed 'alice2.txt'
removed 'alice3.txt'

In [15]: lower = ["foo", "bar", "baz"]

In [16]: upper = ! echo '{"\n".join(lower)}' | tr '[a-z]' '[A-Z]' ⑦
In [17]: upper
Out[17]: ['FOO', 'BAR', 'BAZ']

In [18]: exit
Shutting down kernel
```

- ❶ You can run arbitrary shell commands and pipelines such as `date` or `pip` to install a Python package.

❷

Compare this line of Python code to count the number of lines in *alice.txt* with the invocation of `wc` below it.

- ③ Note that standard output is returned as a list of strings, so to use the value of `total_lines`, you need to get the first item and cast it to an integer.
- ④ Compare this cell and the next to download a file with the invocation of `curl` below it.
- ⑤ You can use Python variables as part of the shell command by using curly braces.
- ⑥ If you want to use literal curly braces, type them twice.
- ⑦ Using a Python variable as standard input can be done, but it gets quite tricky, as you can see.

Jupyter Notebook is, in essence, a browser-based version of Jupyter Console. It supports the same ways to leverage the command line, including the exclamation mark and `bash` magic. The biggest difference is that a notebook can contain not only code but also marked-up text, equations, and data visualizations. It's very popular among data scientists for this reason. Jupyter Notebook is a separate project and environment, but I like to use JupyterLab to work with notebooks, because it offers a more complete IDE.

Figure 10-1 is a screenshot of JupyterLab showing the file explorer (left), a code editor (middle), a notebook (right), and a terminal (bottom). The latter three all show ways to leverage the command line. The code is something I get back to in the next section. This particular notebook is quite similar to the console session I just discussed. The terminal offers a complete shell for you to run command-line tools. Be aware that there's no interactivity possible between this terminal, the code, and the notebook. So this terminal

is not really different from having a separate terminal application open, but it's still helpful when you're working inside a Docker container or on a remote server.

JupyterLab 127.0.0.1:8888/lab

File Edit View Run Kernel Tabs Settings Help

count.py

```
1 #!/usr/bin/env python
2
3 from subprocess import run
4 from sys import argv
5
6 if __name__ == "__main__":
7
8     _, filename, pattern = argv
9
10    with open(filename) as f:
11        alice = f.read()
12
13    words = "\n".join(alice.split())
14
15    grep = run(["grep", "-i", pattern],
16               input=words,
17               capture_output=True,
18               text=True)
19
20    print(len(grep.stdout.strip().split("\n")))
21
```

Untitled1337.ipynb

```
[1]: head alice.txt
Project Gutenberg's Alice's Adventures in Wonderland, by Lewis Carroll

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almost no restrictions whatsoever. You may copy it, give it away or
re-use it under the terms of the Project Gutenberg License included
with this eBook or online at www.gutenberg.org

Title: Alice's Adventures in Wonderland
```

```
[2]: %%bash --out count
grep -oE '\w+' alice.txt |
grep -i alice |
wc -l
```

```
[3]: print(f"Alice appears {int(count)} times in the book")
Alice appears 403 times in the book
```

jovyan@f0c36d91b8a6:~

```
$ head alice.txt
Project Gutenberg's Alice's Adventures in Wonderland, by Lewis Carroll

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almost no restrictions whatsoever. You may copy it, give it away or
re-use it under the terms of the Project Gutenberg License included
with this eBook or online at www.gutenberg.org

Title: Alice's Adventures in Wonderland
```

Saving completed

Figure 10-1. JupyterLab with the file explorer, a code editor, a notebook, and a terminal

WARNING

The notebook in [Figure 10-1](#) also contains a cell using so-called `bash` magic, which allows you to write multiline Bash scripts. Because it's much more difficult to use Python variables, I don't recommend this approach. You're better off creating a Bash script in a separate file and then executing it by using the exclamation mark (!).

Python

The `subprocess` module allows you to run command-line tools from Python and connect to their standard input and output. This module is recommended over the older `os.system()` function. It's not run in a shell by default, but it's possible to change that with the `shell` argument to `run()` function:

```
$ bat count.py
-----
| File: count.py
-----
1 | #!/usr/bin/env python
2 |
3 | from subprocess import run ①
4 | from sys import argv
5 |
6 | if __name__ == "__main__":
7 |
8 |     _, filename, pattern = argv
9 |
10 |    with open(filename) as f: ②
11 |        alice = f.read()
12 |
13 |    words = "\n".join(alice.split()) ③
14 |
15 |    grep = run(["grep", "-i", pattern], ④
16 |               input=words,
17 |               capture_output=True,
18 |               text=True)
```

```
19 |  
20 |     print(len(grep.stdout.strip().split("\n")))
```

- ❶ The recommended way to leverage the command line is to use the `run()` function of the `subprocess` module.
- ❷ Open the file *filename*.
- ❸ Split the entire text into words.
- ❹ Run the command-line tool `grep`, where *words* is passed as standard input.
- ❺ The standard output is available as one long string. Here, I split it on each newline character to count the number of occurrences of *pattern*.

This command-line tool is used as follows:

```
$ ./count.py alice.txt alice  
403
```

Notice that the first argument of the `run` call on line 15 is a list of strings, where the first item is the name of the command-line tool and the remaining items are arguments. This is different from passing a single string. This also means that you don't have any other shell syntax available that would allow for things such as redirection and piping.

R

In R, there are several ways to leverage the command line.

In the following example, I start an R session and count the number of occurrences of the string *alice* in the book *Alice's Adventures in Wonderland* using the `system2()` function:

```
$ R --quiet
> lines <- readLines("alice.txt") ❶
> head(lines)
[1] "Project Gutenberg's Alice's Adventures in Wonderland, by Lewis Carroll"
[2] ""
[3] "This eBook is for the use of anyone anywhere at no cost and with"
[4] "almost no restrictions whatsoever. You may copy it, give it away or"
[5] "re-use it under the terms of the Project Gutenberg License included"
[6] "with this eBook or online at www.gutenberg.org"
> words <- unlist(strsplit(lines, " ")) ❷
> head(words)
[1] "Project"      "Gutenberg's"   "Alice's"       "Adventures"    "in"
[6] "Wonderland,"
> alice <- system2("grep", c("-i", "alice"), input = words, stdout = TRUE) ❸
> head(alice)
[1] "Alice's"      "Alice's"      "ALICE'S"      "ALICE'S"      "Alice"        "Alice"
> length(alice) ❹
[1] 403
```

- ❶ Read in the file *alice.txt*.
- ❷ Split the text into words.
- ❸ Invoke the command-line tool `grep` to keep only the lines that match the string *alice*. The character vector *words* is passed as standard input.
- ❹ Count the number of elements in the character vector *alice*.

A disadvantage of `system2()` is that it first writes the character vector to a file before passing it as standard input to the command-line tool. This can be problematic when dealing with a lot of data and a lot of invocations.

It's better to use a named pipe, because then no data will be written to disk, which is much more efficient. This can be done with the `pipe()` and

`fifo()` functions.¹ The following code demonstrates this:

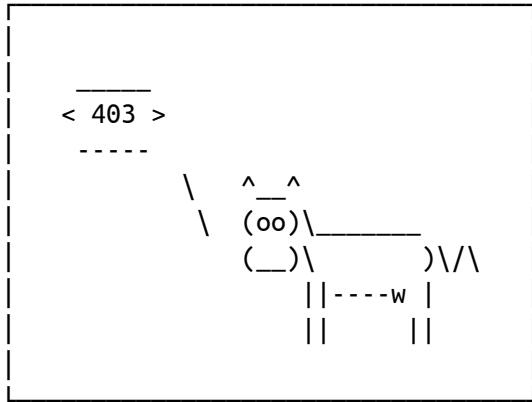
```
> out_con <- fifo("out", "w+") ①
> in_con <- pipe("grep b > out") ②
> writeLines(c("foo", "bar"), in_con) ③
> readLines(out_con) ④
[1] "bar"
> close(out_con); close(in_con); unlink("out") ⑤
```

- ① The function `fifo()` creates a special first-in-first-out file called *out*. This is just a reference to a pipe connection (like standard input and standard output are). No data is actually written to disk.
- ② The tool `grep` will keep only lines that contain a *b* and write them to the named pipe *out*.
- ③ Write two values to standard input of the shell command.
- ④ Read the standard output produced by `grep` as a character vector.
- ⑤ Clean up the connections and delete the special file.

Because this requires quite a bit of boilerplate code (creating connections, writing, reading, cleaning up), I have written a helper function `sh()`. Using the pipe operator (`%>%`) from the `magrittr` package, I chain together multiple shell commands:

```
> library(magrittr)
>
> sh <- function(.data, command) {
+   temp_file <- tempfile()
+   out_con <- fifo(temp_file, "w+")
+   in_con <- pipe(paste0(command, " > ", temp_file))
+   writeLines(as.character(.data), in_con)
+   result <- readLines(out_con)
+   close(out_con)
+   close(in_con)
```

```
+   unlink(temp_file)
+   result
+ }
>
> lines <- readLines("alice.txt")
> words <- unlist(strsplit(lines, " "))
>
> sh(words, "grep -i alice") %>%
+   sh("wc -l") %>%
+   sh("cowsay") %>%
+   cli::cat_boxx()
```



```
>
> q("no")
```

RStudio

The RStudio IDE is arguably the most popular environment for working with R. When you open RStudio, you first see the console tab, as shown in Figure 10-2.

2e - master - RStudio

The screenshot shows the RStudio interface with the following components:

- Code Editor:** scratch.R contains R code for reading lines from a file, piping them through sh, wc, and cowsay, and then piping the result to cat_boxx.
- Environment:** The Global Environment pane shows two objects: "lines" (character [1:3735] "Project Gutenberg's Alice's Adventu...") and "words" (Large character (30754 elements, 599.1 kB)).
- Help:** The Help pane is titled "connections {base}" and describes functions to manipulate connections (files, URLs, etc.).
- Terminal:** The terminal shows the command `sh("cowsay") %>% cli::cat_boxx()` being run, resulting in the output of the cowsay command.

Functions to Manipulate Connections (Files, URLs, ...)

Description

Functions to create, open and close connections, i.e., “generalized files”, such as possibly compressed files, URLs, pipes, etc.

Usage

```

file(description = "", open = "", blocking = TRUE,
     encoding = getOption("encoding"), raw = FALSE,
     method = getOption("url.method", "default"))

url(description, open = "", blocking = TRUE,
     encoding = getOption("encoding"),
     method = getOption("url.method", "default"),
     headers = NULL)

gzfile(description, open = "", encoding = getOption("encoding"),
       compression = 6)

bzfile(description, open = "", encoding = getOption("encoding"),
       compression = 9)

xzfile(description, open = "", encoding = getOption("encoding"),
       compression = 6)

unz(description, filename, open = "", encoding = getOption("encoding"))

pipe(description, open = "", encoding = getOption("encoding"))

fifo(description, open = "", blocking = FALSE,
      encoding = getOption("encoding"))
  
```

Figure 10-2. RStudio IDE with console tab open

The terminal tab is right next to the console tab. It offers a complete shell, as shown in [Figure 10-3](#).

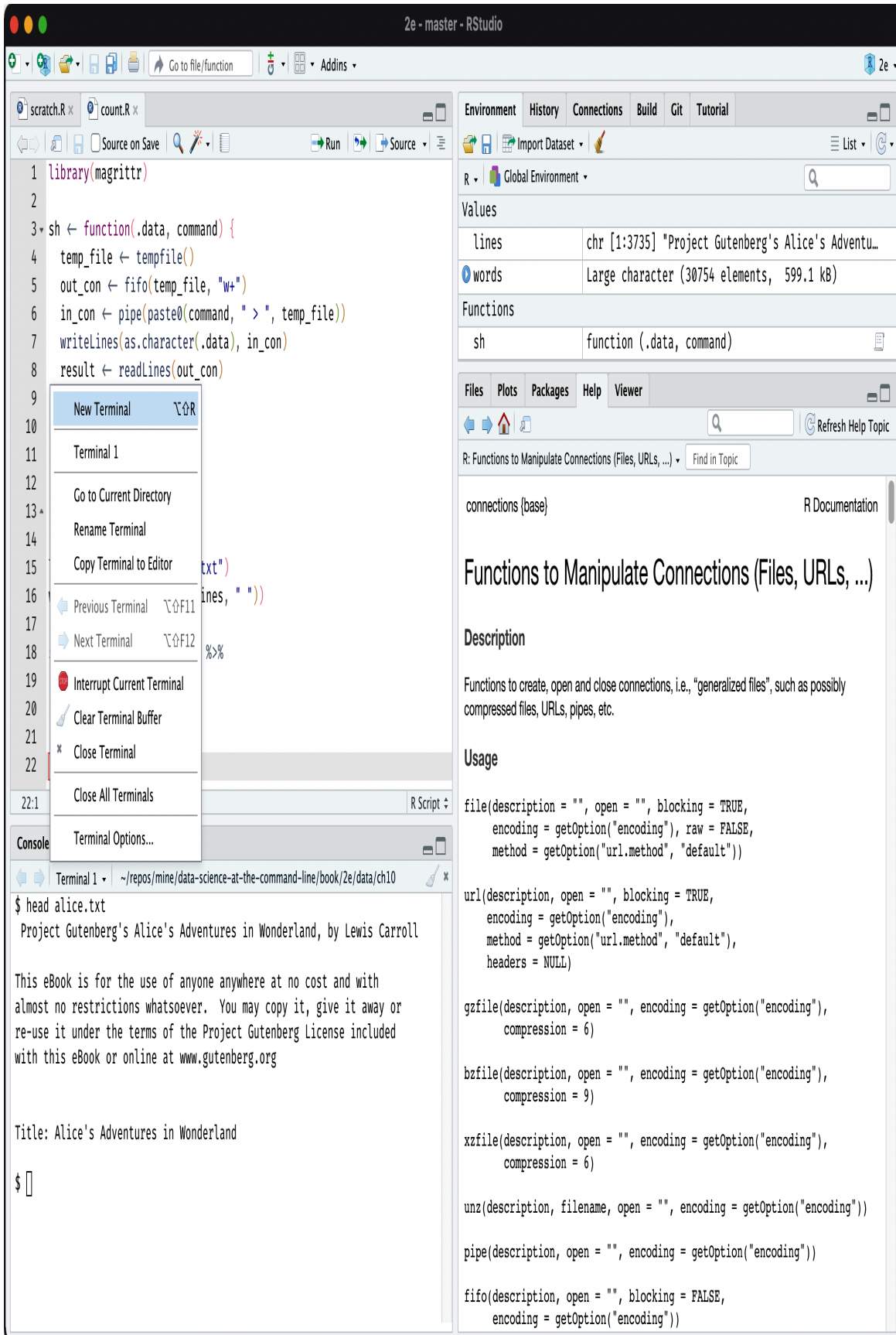


Figure 10-3. RStudio IDE with terminal tab open

Note that, just as with JupyterLab, this terminal is not connected to the console or to any R scripts.

Apache Spark

Apache Spark is a cluster-computing framework. It's the eight-hundred-pound gorilla you turn to when it's impossible to fit your data in memory. Spark itself is written in Scala, but you can also interact with it from Python using [PySpark](#) and from R using [SparkR](#) or [sparklyr](#).

Data processing and machine learning pipelines are defined through a series of transformations and one final action. One of these transformations is the `pipe()` transformation, which allows you to run the entire dataset through a shell command such as a Bash or Perl script. The items in the dataset are written to standard input, and the standard output is returned as a resilient distributed dataset (RDD) of strings.

In the following session, I start a Spark shell and again count the number of occurrences of *alice* in the book *Alice's Adventures in Wonderland*:

```
$ spark-shell --master local[6]
Spark context Web UI available at http://3d1bec8f2543:4040
Spark context available as 'sc' (master = local[6], app id = local-16193763).
Spark session available as 'spark'.
Welcome to
```

```
   _/\_/_\_\_/\_/\_/\_
  / \ \_ \ \_ \_ ` / \_ / \_/
  / \_ / . \_ / \_, \_ / \_ / \_ / \_ \_ version 3.1.1
  / \_ /
```

```
Using Scala version 2.12.10 (OpenJDK 64-Bit Server VM, Java 11.0.10)
Type in expressions to have them evaluated.
Type :help for more information.
```

```
scala> val lines = sc.textFile("alice.txt") ❶
lines: org.apache.spark.rdd.RDD[String] = alice.txt MapPartitionsRDD[1] at
textF
ile at <console>:24
```

```

scala> lines.first()
res0: String = Project Gutenberg's Alice's Adventures in Wonderland, by Lewis
Ca
rroll

scala> val words = lines.flatMap(line => line.split(" ")) ②
words: org.apache.spark.rdd.RDD[String] = MapPartitionsRDD[2] at flatMap at
<con
sole>:25

scala> words.take(5)
res1: Array[String] = Array(Project, Gutenberg's, Alice's, Adventures, in)

scala> val alice = words.pipe("grep -i alice") ③
alice: org.apache.spark.rdd.RDD[String] = PipedRDD[3] at pipe at <console>:25

scala> alice.take(5)
res2: Array[String] = Array(Alice's, Alice's, ALICE'S, ALICE'S, Alice)

scala> val counts = alice.pipe("wc -l") ④
counts: org.apache.spark.rdd.RDD[String] = PipedRDD[4] at pipe at <console>:25

scala> counts.collect()
res3: Array[String] = Array(64, 72, 94, 93, 67, 13) ⑤

scala> counts.map(x => x.toInt).reduce(_ + _) ⑥
res4: Int = 403

scala> sc.textFile("alice.txt").flatMap(line => line.split(" ")).pipe("grep -i
a
lice").pipe("wc -l").map(x => x.toInt).reduce(_ + _)
res5: Int = 403 ⑦

```

- ➊ Read *alice.txt* such that each line is an element.
- ➋ Split each element on spaces. In other words, each line is split into words.
- ➌ Pipe each partition through `grep` to keep only the elements that match the string *alice*.

- ④ Pipe each partition through `wc` to count the number of elements.
- ⑤ There's one count for each partition.
- ⑥ Sum all counts to get a final count. Note that elements first need to be converted from strings to integers.
- ⑦ This shows the preceding steps combined into a single command.

TIP

The `pipe()` transformation is also available in PySpark, SparkR, and sparklyr.

If you want to use a custom command-line tool in your pipeline, then you need to make sure that it's present on all nodes in the cluster (known as the executors). One way to do this is to specify the filename(s) with the `--files` option when you're submitting Spark applications using `spark-submit`.

Bill Chambers and Matei Zaharia (the original authors of Apache Spark) mention in their book *Spark: The Definitive Guide* (O'Reilly) that “[t]he `pipe` method is probably one of Spark's more interesting methods.” That's quite the compliment! I think it's fantastic that the developers of Apache Spark added the ability to leverage a 50-year-old technology.

Summary

In this chapter you learned several ways to leverage the command line in other situations, including programming languages and other environments. It's important to realize that the command line doesn't exist in a vacuum. What matters most is that you use tools, sometimes in combination with one another, that reliably get the job done.

Now that we've had the four OSEMN chapters and the four intermezzo chapters, it's time to wrap this up in the final chapter.

For Further Exploration

There are also ways to integrate two programming languages directly without the use of the command line. For example, the [reticulate package](#) in R allows you to interface with Python directly.

¹ Thanks to Jim Hester for suggesting this.

Chapter 11. Conclusion

In this final chapter, the book comes to a close. I'll first recap what I've discussed in the previous 10 chapters, and then I'll offer you three pieces of advice and provide some resources to further explore the related topics we touched on. Finally, in case you have any questions, comments, or new command-line tools to share, I provide a few ways to get in touch with me.

Let's Recap

This book explored the power of using the command line to do data science. I find it an interesting observation that the challenges posed by this relatively young field can be tackled using such a time-tested technology. I hope that you now see what the command line is capable of. The many command-line tools offer all sorts of possibilities that are well suited to the variety of tasks encompassing data science.

There are many definitions of data science available. In [Chapter 1](#), I introduced the five-step OSEMN model as defined by Hilary Mason and Chris Wiggins, because it is a very practical one that translates to very specific tasks. The acronym OSEMN stands for *obtaining, scrubbing, exploring, modeling, and interpreting* data. [Chapter 1](#) also explained why the command line is very suitable for doing these data science tasks.

In [Chapter 2](#), I explained how you can get all the tools used in this book. [Chapter 2](#) also provided an introduction to the essential tools and concepts of the command line.

The four OSEMN model chapters focused on performing those practical tasks using the command line. I did not devote a chapter to the fifth step, interpreting data, because quite frankly, the computer—let alone the command line—is of very little use here. I have, however, provided some pointers for further reading on this topic.

In the four intermezzo chapters, we looked at some broader topics of doing data science at the command line, topics that are not really specific to a particular step. In [Chapter 4](#), I explained how you can turn one-liners and existing code into reusable command-line tools. In [Chapter 6](#), I described how you can manage your data workflow using a tool called `make`. In [Chapter 8](#), I demonstrated how ordinary command-line tools and pipelines can be run in parallel using GNU Parallel. In [Chapter 10](#), I showed that the command line doesn't exist in a vacuum but can be leveraged from other programming languages and environments. The topics discussed in these intermezzo chapters can be applied at any point in your data workflow.

It's impossible to demonstrate all the command-line tools that are available and relevant for doing data science. New tools are created on a daily basis. As you may have come to understand by now, this book is more about the idea of using the command line rather than giving you an exhaustive list of tools.

Three Pieces of Advice

You probably spent a good bit of time reading these chapters and perhaps also following along with the code examples. In the hope that it maximizes the return on this investment and increases the probability that you'll continue to incorporate the command line into your data science workflow, I would like to offer you three pieces of advice: (1) be patient, (2) be creative, and (3) be practical. In the next three subsections I elaborate on each piece of advice.

Be Patient

The first piece of advice that I can give is to *be patient*. Working with data on the command line is different from using a programming language, and therefore it requires a different mindset.

Moreover, the command-line tools themselves are not without their quirks and inconsistencies. This is partly because they have been developed by

many different people over the course of multiple decades. If you ever find yourself at a loss regarding their mind-dazzling options, don't forget to use `--help`, `man`, `tldr`, or your favorite search engine to learn more.

Still, especially in the beginning, working on the command line can be a frustrating experience. Trust me, you'll become more proficient as you practice using the command line and its tools. The command line has been around for many decades and will be around for many more to come. It's a worthwhile investment.

Be Creative

The second, related piece of advice is to *be creative*. The command line is very flexible. By combining the command-line tools, you can accomplish more than you might think.

I encourage you to not immediately fall back onto your programming language. And when you do have to use a programming language, think about whether the code can be generalized or reused in some way; if so, consider creating your own command-line tool with that code using the steps I discussed in [Chapter 4](#). If you believe your tool may be beneficial for others, you could even go one step further by making it open source. Maybe there's a step you know how to perform at the command line, but you would rather not leave the comfort of the main programming language or environment you're working in. Perhaps you can use one of the approaches listed in [Chapter 10](#).

Be Practical

The third piece of advice is to *be practical*. Being practical is related to being creative but deserves a separate explanation. In the previous subsection, I mentioned that you should not immediately fall back to a programming language. Of course, the command line has its limits. Throughout the book, I have emphasized that the command line should be regarded as a companion approach to doing data science.

I've discussed four steps for doing data science at the command line. In practice, the applicability of the command line is higher for step 1 than it is for step 4. You should use whichever approach works best for the task at hand. And it's perfectly fine to mix and match approaches at any point in your workflow. As I've shown in [Chapter 10](#), the command line is wonderful at being integrated with other approaches, programming languages, and statistical environments. There's a certain trade-off with each approach, and part of becoming proficient at the command line is to learn when to use which approach.

In conclusion, when you're patient, creative, and practical, the command line will make you a more efficient and productive data scientist.

Where to Go from Here

As this book is on the intersection of the command line and data science, many related topics have only been touched on. Now it's up to you to further explore these topics. The following subsections provide a list of topics and suggested resources to consult.

The Command Line

- *The Linux Command Line: A Complete Introduction*, 2nd ed., by William Shotts (San Francisco: No Starch, 2019)
- *Unix Power Tools*, 3rd ed., by Shelley Powers, Jerry Peek, Tim O'Reilly, and Mike Loukides (Sebastopol, CA: O'Reilly, 2002)
- *Learning the vi and Vim Editors*, 7th ed., by Arnold Robbins, Elbert Hannah, and Linda Lamb (Sebastopol, CA: O'Reilly, 2008)

Shell Programming

- *Classic Shell Scripting* by Arnold Robbins and Nelson H. F. Beebe (Sebastopol, CA: O'Reilly, 2005)

- *Wicked Cool Shell Scripts*, 2nd ed., by Dave Taylor and Brandon Perry (San Franciscio: No Starch, 2017)
- *bash Cookbook*, 2nd ed., by Carl Albing and JP Vossen (Sebastopol, CA: O'Reilly, 2017)

Python, R, and SQL

- *Learn Python 3 the Hard Way* by Zed A. Shaw (Boston: Addison-Wesley, 2017)
- *Python for Data Analysis*, 2nd ed., by Wes McKinney (Sebastopol, CA: O'Reilly, 2017)
- *Data Science from Scratch*, 2nd ed., by Joel Grus (Sebastopol, CA: O'Reilly, 2019)
- *R for Data Science* by Hadley Wickham and Garrett Grolemund (Sebastopol, CA: O'Reilly, 2016)
- *R for Everyone*, 2nd ed., by Jared P. Lander (Boston: Addison-Wesley, 2017)
- *Sams Teach Yourself SQL in 10 Minutes a Day*, 5th ed., by Ben Forta (Hoboken: Sams, 2020)

APIs

- *Mining the Social Web*, 3rd ed., by Matthew A. Russell and Mikhail Klassen (Sebastopol, CA: O'Reilly, 2019)
- *Data Source Handbook* by Pete Warden (Sebastopol, CA: O'Reilly, 2011)

Machine Learning

- *Python Machine Learning*, 3rd ed., by Sebastian Raschka and Vahid Mirjalili (Birmingham, UK: Packt, 2019)

- *Pattern Recognition and Machine Learning* by Christopher M. Bishop (New York: Springer, 2006)
- *Information Theory, Inference, and Learning Algorithms* by David J. C. MacKay (Cambridge, UK: Cambridge University Press, 2003)

Getting in Touch

This book would not have been possible without the many people who created the command line and the numerous tools. It's safe to say that the current ecosystem of command-line tools for data science is a community effort. I have only been able to give you a glimpse of the many command-line tools available. New ones are created every day, and perhaps some day you will create one yourself, in which case, I would love to hear from you. I'd also appreciate it if you would drop me a line whenever you have a question, comment, or suggestion. There are a few ways to get in touch:

- Email: jeroen@jeroenjanssens.com
- Twitter: [@jeroenhjanssens](https://twitter.com/jeroenhjanssens)
- Book website: <https://datascienceatthecli.com>
- Book GitHub repository: <https://github.com/jeroenjanssens/datascience-at-the-command-line>

Thank you.

Appendix. List of Command-Line Tools

This is an overview of all the command-line tools discussed in this book. This includes binary executables, interpreted scripts, and Z Shell builtins and keywords. For each command-line tool, the following information, when available and appropriate, is provided:

- The actual command to type at the command line
- A description
- The version used in the book
- The year that version was released
- The primary author(s)
- A website to find more information
- How to obtain help
- An example usage

All command-line tools listed here are included in the Docker image. See [Chapter 2](#) for instructions on how to set it up. Please note that citing open source software is not trivial, and that some information may be missing or incorrect.

alias

Define or display aliases. `alias` is a Z shell builtin.

```
$ type alias
alias is a shell builtin
```

```
$ man zshbuiltins | grep -A 10 alias

$ alias l
l='ls --color -lhF --group-directories-first'

$ alias python=python3
```

awk

Pattern scanning and text processing language. **awk** (version 1.3.4) by Mike D. Brennan and Thomas E. Dickey (2019). More information: <https://invisible-island.net/mawk>.

```
$ type awk
awk is /usr/bin/awk

$ man awk

$ seq 5 | awk '{sum+=$1} END {print sum}'
15
```

aws

Unified tool to manage AWS services. **aws** (version 2.1.32) by Amazon Web Services (2021). More information: <https://aws.amazon.com/cli>.

```
$ type aws
aws is /usr/local/bin/aws

$ aws --help
```

bash

GNU Bourne-Again Shell. **bash** (version 5.0.17) by Brian Fox and Chet Ramey (2019). More information: <https://www.gnu.org/software/bash>.

```
$ type bash
bash is /usr/bin/bash
```

```
$ man bash
```

bat

A `cat` clone with syntax highlighting and Git integration. `bat` (version 0.18.0) by David Peter (2021). More information: <https://github.com/sharkdp/bat>.

```
$ type bat
bat is an alias for bat --tabs 8 --paging never

$ man bat
```

bc

An arbitrary precision calculator language. `bc` (version 1.07.1) by Philip A. Nelson (2017). More information: <https://www.gnu.org/software/bc>.

```
$ type bc
bc is /usr/bin/bc

$ man bc

$ bc -l <<< 'e(1)'
2.71828182845904523536
```

body

Apply expression to all but the first line. `body` (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type body
body is /usr/bin/dsutils/body

$ seq 10 | header -a 'values' | body shuf
values
```

```
7  
5  
2  
6  
10  
3  
9  
8  
1  
4
```

cat

Concatenate files and print on the standard output. **cat** (version 8.30) by Torbjorn Granlund and Richard M. Stallman (2018). More information: <https://www.gnu.org/software/coreutils>.

```
$ type cat  
cat is /usr/bin/cat  
  
$ man cat  
  
$ cat *.log > all.log
```

cd

Change the shell working directory. **cd** is a Z shell builtin.

```
$ type cd  
cd is a shell builtin  
  
$ man zshbuiltins | grep -A 10 cd  
  
$ cd ~  
  
$ pwd  
/home/dst  
  
$ cd ..  
  
$ pwd  
/home
```

```
$ cd /data/ch01
```

chmod

Change file mode bits. `chmod` (version 8.30) by David MacKenzie and Jim Meyering (2018). I use `chmod` in [Chapter 4](#) to make a tool executable. More information: <https://www.gnu.org/software/coreutils>.

```
$ type chmod
chmod is /usr/bin/chmod

$ man chmod

$ chmod u+x script.sh
```

cols

Apply command to a subset of columns. `cols` (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type cols
cols is /usr/bin/dsutils/cols
```

column

Columnate lists. `column` (version 2.36.1) by Karel Zak (2021). More information: <https://www.kernel.org/pub/linux/utils/util-linux>.

```
$ type column
column is /usr/bin/column
```

cowsay

Configurable speaking cow. **cowsay** (version 3.0.3) by Tony Monroe (1999). More information: <https://github.com/tnalpgge/rank-amateur-cowsay>.

```
$ type cowsay
cowsay is /usr/bin/cowsay

$ man cowsay

$ echo 'The command line is awesome!' | cowsay
< The command line is awesome! >
-----
 \  ^__^
  \  (oo)\_____
    (__)\       )\/\
        ||----w |
        ||     ||
```

cp

Copy files and directories. **cp** (version 8.30) by Torbjorn Granlund, David MacKenzie, and Jim Meyering (2018). More information: <https://www.gnu.org/software/coreutils>.

```
$ type cp
cp is /usr/bin/cp

$ man cp

$ cp -r ~/Downloads/*.xlsx /data
```

csv2vw

Convert CSV to Vowpal Wabbit format. **csv2vw** (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type csv2vw  
csv2vw is /usr/bin/dsutils/csv2vw
```

csvcut

Filter and truncate CSV files. `csvcut` (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvcut  
csvcut is /usr/bin/csvcut  
  
$ csvcut --help  
  
$ csvcut -c bill,tip /data/ch05/tips.csv | trim  
bill,tip  
16.99,1.01  
10.34,1.66  
21.01,3.5  
23.68,3.31  
24.59,3.61  
25.29,4.71  
8.77,2.0  
26.88,3.12  
15.04,1.96  
... with 235 more lines
```

csvgrep

Search CSV files. `csvgrep` (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvgrep  
csvgrep is /usr/bin/csvgrep  
  
$ csvgrep --help
```

csvjoin

Execute a SQL-like join to merge CSV files on a specified column or columns. `csvjoin` (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvjoin
csvjoin is /usr/bin/csvjoin

$ csvjoin --help
```

csvlook

Render a CSV file in the console as a Markdown-compatible, fixed-width table. `csvlook` (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvlook
csvlook is a shell function

$ csvlook --help

$ csvlook /data/ch05/tips.csv
+-----+-----+-----+-----+-----+-----+-----+
| bill | tip | sex | smoker | day | time | size |
+-----+-----+-----+-----+-----+-----+-----+
| 16.99 | 1.01 | Female | False | Sun | Dinner | 2 |
| 10.34 | 1.66 | Male | False | Sun | Dinner | 3 |
| 21.01 | 3.50 | Male | False | Sun | Dinner | 3 |
| 23.68 | 3.31 | Male | False | Sun | Dinner | 2 |
| 24.59 | 3.61 | Female | False | Sun | Dinner | 4 |
| 25.29 | 4.71 | Male | False | Sun | Dinner | 4 |
| 8.77 | 2.00 | Male | False | Sun | Dinner | 2 |
| 26.88 | 3.12 | Male | False | Sun | Dinner | 4 |
... with 236 more lines
```

csvquote

Enable common Unix utilities to work correctly with CSV data. `csvquote` (version 0.1) by Dan Brown (2018). More information:

<https://github.com/dbro/csvquote>.

```
$ type csvquote
csvquote is /usr/local/bin/csvquote
```

csvsort

Sort CSV files. **csvsort** (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvsort
csvsort is /usr/bin/csvsort

$ csvsort --help
```

csvsql

Execute SQL statements on CSV files. **csvsql** (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvsql
csvsql is /usr/bin/csvsql

$ csvsql --help
```

csvstack

Stack up the rows from multiple CSV files. **csvstack** (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvstack
csvstack is /usr/bin/csvstack

$ csvstack --help
```

csvstat

Print descriptive statistics for each column in a CSV file. **csvstat** (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type csvstat
csvstat is /usr/bin/csvstat

$ csvstat --help
```

curl

Transfer a URL. **curl** (version 7.68.0) by Daniel Stenberg (2016). More information: <https://curl.haxx.se>.

```
$ type curl
curl is /usr/bin/curl

$ man curl
```

cut

Remove sections from each line of files. **cut** (version 8.30) by David M. Ihnat, David MacKenzie, and Jim Meyering (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type cut
cut is /usr/bin/cut

$ man cut
```

display

Display an image or image sequence on any X server. **display** (version 6.9.10-23) by ImageMagick Studio LLC (2019). More information: <https://imagemagick.org>.

```
$ type display  
display is a shell function
```

dseq

Generate a sequence of dates. `dseq` (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type dseq  
dseq is /usr/bin/dsutils/dseq  
  
$ dseq 3  
2021-06-30  
2021-07-01  
2021-07-02
```

echo

Display a line of text. `echo` (version 8.30) by Brian Fox and Chet Ramey (2019). Useful for using literal text as standard input to the next tool. More information: <https://www.gnu.org/software/coreutils>.

```
$ type echo  
echo is a shell builtin  
  
$ man echo  
  
$ echo Hippopotomonstrosesquippedaliophobia | cowsay  
-----  
< Hippopotomonstrosesquippedaliophobia >  
-----  
 \ ^__^  
  \  (oo)\_____  
   (__)\       )\/\|  
    ||----w |  
    ||     ||  
  
$ echo -n Hippopotomonstrosesquippedaliophobia | wc -c  
36
```

env

Run a program in a modified environment. `env` (version 8.32) by Richard Mlynarik, David MacKenzie, and Assaf Gordon (2020). More information: <https://www.gnu.org/software/coreutils>.

```
$ type env  
env is /usr/bin/env  
  
$ man env
```

export

Set export attribute for shell variables. Useful for making shell variables available to other command-line tools. `export` is a Z shell builtin.

```
$ type export  
export is a reserved word  
  
$ man zshbuiltins | grep -A 10 export  
  
$ export PATH="$PATH:$HOME/bin"
```

fc

Control the interactive history mechanism. `fc` is a Z shell builtin. I use `fc` in [Chapter 4](#) to edit the command in `nano`.

```
$ type fc  
fc is a shell builtin  
  
$ man zshbuiltins | grep -A 10 '^ *fc '
```

find

Search for files in a directory hierarchy. `find` (version 4.7.0) by Eric B. Decker, James Youngman, and Kevin Dalley (2019). More information:

<https://www.gnu.org/software/findutils>.

```
$ type find
find is /usr/bin/find

$ man find

$ find /data -type f -name '*.csv' -size -3
/data/ch03/tmnt-basic.csv
/data/ch03/tmnt-missing-newline.csv
/data/ch03/tmnt-with-header.csv
/data/ch05/names-comma.csv
/data/ch05/irismeta.csv
/data/ch05/names.csv
/data/ch07/datatypes.csv
```

fold

Wrap each input line to fit in specified width. **fold** (version 8.30) by David MacKenzie (2020). More information:

<https://www.gnu.org/software/coreutils>.

```
$ type fold
fold is /usr/bin/fold

$ man fold
```

for

Execute commands for each member in a list. **for** is a Z shell builtin. In [Chapter 8](#), I discuss the advantages of using **parallel** instead of **for**.

```
$ type for
for is a reserved word

$ man zshmisc | grep -EA 10 '^ *for '
$ for i in {A..C} "It's easy as" {1..3}; do echo $i; done
A
B
C
```

```
It's easy as
1
2
3
```

fx

Interactive JSON viewer. `fx` (version 20.0.2) by Anton Medvedev (2020). More information: <https://github.com/antonmedv/fx>.

```
$ type fx
fx is /usr/local/bin/fx

$ fx --help

$ echo '[1,2,3]' | fx 'this.map(x => x * 2)'
[
  2,
  4,
  6
]
```

git

The stupid content tracker. `git` (version 2.25.1) by Linus Torvalds and Junio C. Hamano (2021). More information: <https://git-scm.com>.

```
$ type git
git is /usr/bin/git

$ man git
```

grep

Print lines that match patterns. `grep` (version 3.4) by Jim Meyering (2019). More information: <https://www.gnu.org/software/grep>.

```
$ type grep
grep is /usr/bin/grep
```

```
$ man grep  
  
$ seq 100 | grep 3 | wc -l  
19
```

gron

Make JSON greppable. **gron** (version 0.6.1) by Tom Hudson (2021). More information: <https://github.com/TomNomNom/gron>.

```
$ type gron  
gron is /usr/bin/gron  
  
$ man gron
```

head

Output the first part of files. **head** (version 8.30) by David MacKenzie and Jim Meyering (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type head  
head is /usr/bin/head  
  
$ man head  
  
$ seq 100 | head -n 5  
1  
2  
3  
4  
5
```

header

Add, replace, and delete header lines. **header** (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type header  
header is /usr/bin/dsutils/header
```

history

GNU History Library. **history** (version 8.1) by Brian Fox and Chet Ramey (2020). More information: <https://www.gnu.org/software/bash>.

```
$ type history  
history is a shell builtin
```

hostname

Show or set the system's host name. **hostname** (version 3.23) by Peter Tobias, Bernd Eckenfels, and Michael Meskes (2021). More information: <https://sourceforge.net/projects/net-tools/>.

```
$ type hostname  
hostname is /usr/bin/hostname  
  
$ man hostname  
  
$ hostname  
faa90036a8ee  
  
$ hostname -i  
172.17.0.2
```

in2csv

Convert common, but less awesome, tabular data formats to CSV. **in2csv** (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type in2csv  
in2csv is /usr/bin/in2csv  
  
$ in2csv --help
```

jq

Command-line JSON processor. jq (version 1.6) by Stephen Dolan (2021). More information: <https://stedolan.github.io/jq>.

```
$ type jq
jq is /usr/bin/jq

$ man jq
```

json2csv

Convert JSON to CSV. json2csv (version 1.2.1) by Jehiah Czebotar (2019). More information: <https://github.com/jehiah/json2csv>.

```
$ type json2csv
json2csv is /usr/bin/json2csv

$ json2csv --help
```

|
List directory contents in long format with directories grouped before files, human-readable file sizes, and access permissions. l by Unknown (1999).

```
$ type l
l is an alias for ls --color -lhF --group-directories-first

$ cd /data/ch03

$ ls
logs.tar.gz      tmnt-basic.csv          tmnt-with-header.csv
r-datasets.db    tmnt-missing-newline.csv top2000.xlsx

$ l
total 924K
-rw-r--r-- 1 dst dst 627K Jun 29 14:36 logs.tar.gz
-rw-r--r-- 1 dst dst 189K Jun 29 14:36 r-datasets.db
-rw-r--r-- 1 dst dst 149 Jun 29 14:36 tmnt-basic.csv
-rw-r--r-- 1 dst dst 148 Jun 29 14:36 tmnt-missing-newline.csv
```

```
-rw-r--r-- 1 dst dst 181 Jun 29 14:36 tmnt-with-header.csv  
-rw-r--r-- 1 dst dst 91K Jun 29 14:36 top2000.xlsx
```

less

Opposite of more. **less** (version 551) by Mark Nudelman (2019). More information: <https://www.greenwoodsoftware.com/less>.

```
$ type less  
less is an alias for less -R  
  
$ man less  
  
$ less README
```

ls

List directory contents. **ls** (version 8.30) by Richard M. Stallman and David MacKenzie (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type ls  
ls is /usr/bin/ls  
  
$ man ls  
  
$ ls /data  
ch01 ch02 ch03 ch04 ch05 ch06 ch07 ch08 ch09 ch10
```

make

A program for maintaining computer programs. **make** (version 4.3) by Stuart I. Feldman (2020). More information: <https://www.gnu.org/software/make>.

```
$ type make  
make is /usr/bin/make
```

```
$ man make
```

```
$ make sandwich
```

man

An interface to the system reference manuals. `man` (version 2.9.1) by John W. Eaton and Colin Watson (2020). More information:

<https://nongnu.org/man-db>.

```
$ type man
man is /usr/bin/man
```

```
$ man man
```

```
$ man excel
No manual entry for excel
```

mkdir

Make directories. `mkdir` (version 8.30) by David MacKenzie (2019). More information: *<https://www.gnu.org/software/coreutils>*.

```
$ type mkdir
mkdir is /usr/bin/mkdir
```

```
$ man mkdir
```

```
$ mkdir -p /data/ch{01..10}
```

mv

Move (rename) files. `mv` (version 8.30) by Mike Parker, David MacKenzie, and Jim Meyering (2020). More information:

<https://www.gnu.org/software/coreutils>.

```
$ type mv
mv is /usr/bin/mv
```

```
$ man mv  
$ mv results{,.bak}
```

nano

Nano's ANOther editor, inspired by Pico. `nano` (version 5.4) by Benno Schulenberg, David Lawrence Ramsey, Jordi Mallach, Chris Allegretta, Robert Siemborski, and Adam Rogoyski (2020). More information:
<https://nano-editor.org>.

```
$ type nano  
nano is /usr/bin/nano
```

nl

Number lines of files. `nl` (version 8.30) by Scott Bartram and David MacKenzie (2020). More information:
<https://www.gnu.org/software/coreutils>.

```
$ type nl  
nl is /usr/bin/nl  
  
$ man nl  
  
$ nl /data/ch05/alice.txt | head  
1 Project Gutenberg's Alice's Adventures in Wonderland, by Lewis Carroll  
2  
3 This eBook is for the use of anyone anywhere at no cost and with  
4 almost no restrictions whatsoever. You may copy it, give it away or  
5 re-use it under the terms of the Project Gutenberg License included  
6 with this eBook or online at www.gutenberg.org  
7  
8  
9 Title: Alice's Adventures in Wonderland  
10
```

parallel

Build and execute shell command lines from standard input in parallel.

`parallel` (version 20161222) by Ole Tange (2016). More information:

<https://www.gnu.org/software/parallel>.

```
$ type parallel
parallel is /usr/bin/parallel

$ man parallel

$ seq 3 | parallel "echo Processing file {} .csv"
Processing file 1.csv
Processing file 2.csv
Processing file 3.csv
```

paste

Merge lines of files. `paste` (version 8.30) by David M. Ihnat and David

MacKenzie (2019). More information:

<https://www.gnu.org/software/coreutils>.

```
$ type paste
paste is /usr/bin/paste

$ man paste

$ paste -d, <(seq 5) <(dseq 5)
1,2021-06-30
2,2021-07-01
3,2021-07-02
4,2021-07-03
5,2021-07-04

$ seq 5 | paste -sd+
1+2+3+4+5
```

pbc

Parallel bc. pbc (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type pbc
pbc is /usr/bin/dsutils/pbc

$ seq 3 | pbc '{1}^2'
1
4
9
```

pip

A tool for installing and managing Python packages. pip (version 20.0.2) by PyPA (2020). More information: <https://pip.pypa.io>.

```
$ type pip
pip is /usr/bin/pip

$ man pip

$ pip install pandas

$ pip freeze | grep sci
scikit-learn==0.24.2
scipy==1.7.0
```

pup

Parsing HTML at the command line. pup (version 0.4.0) by Eric Chiang (2016). More information: <https://github.com/EricChiang/pup>.

```
$ type pup
pup is /usr/bin/pup

$ pup --help
```

pwd

Print name of current/working directory. **pwd** (version 8.30) by Jim Meyering (2019). More information:
<https://www.gnu.org/software/coreutils>.

```
$ type pwd  
pwd is a shell builtin  
  
$ man pwd  
  
$ cd ~  
  
$ pwd  
/home/dst
```

python

An interpreted, interactive, object-oriented programming language. **python** (version 3.8.5) by the Python Software Foundation (2021). More information: *<https://www.python.org>.*

```
$ type python  
python is an alias for python3  
  
$ man python
```

R

A language and environment for statistical computing. **R** (version 4.0.4) by the R Foundation for Statistical Computing (2021). More information: *<https://www.r-project.org>.*

```
$ type R  
R is /usr/bin/R  
  
$ man R
```

rev

Reverse lines characterwise. `rev` (version 2.36.1) by Karel Zak (2021). More information: <https://www.kernel.org/pub/linux/utils/util-linux>.

```
$ type rev
rev is /usr/bin/rev

$ echo 'Satire: Veritas' | rev
satireV :eritaS

$ echo 'Ça va?' | rev | cut -c 2- | rev
Ça va
```

rm

Remove files or directories. `rm` (version 8.30) by Paul Rubin, David MacKenzie, Richard M. Stallman, and Jim Meyering (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type rm
rm is /usr/bin/rm

$ man rm

$ rm *.old
```

rush

R One-Liners from the Shell. `rush` (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/rush>.

```
$ type rush
rush is /usr/local/lib/R/site-library/rush/exec/rush

$ rush --help

$ rush run '6*7'
```

```
$ rush run --tidyverse 'filter(starwars, species == "Human") %>% select(name)'  
# A tibble: 35 x 1  
  name  
  <chr>  
1 Luke Skywalker  
2 Darth Vader  
3 Leia Organa  
4 Owen Lars  
5 Beru Whitesun lars  
6 Biggs Darklighter  
7 Obi-Wan Kenobi  
8 Anakin Skywalker  
9 Wilhuff Tarkin  
10 Han Solo  
# ... with 25 more rows
```

sample

Filter lines from standard input according to some probability, with a given delay, and for a certain duration. **sample** (version 0.2.4) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/sample>.

```
$ type sample  
sample is /usr/local/bin/sample  
  
$ sample --help  
  
$ seq 1000 | sample -r 0.01 | trim 5  
481  
503  
695  
940
```

scp

OpenSSH secure file copy. **scp** (version 1:8.2p1-4ubuntu0.2) by Timo Rinne and Tatu Ylonen (2019). More information: <https://www.openssh.com>.

```
$ type scp  
scp is /usr/bin/scp
```

```
$ man scp
```

sed

Stream editor for filtering and transforming text. **sed** (version 4.7) by Jay Fenlason, Tom Lord, Ken Pizzini, and Paolo Bonzini (2018). More information: <https://www.gnu.org/software/sed>.

```
$ type sed
sed is /usr/bin/sed
```

```
$ man sed
```

seq

Print a sequence of numbers. **seq** (version 8.30) by Ulrich Drepper (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type seq
seq is /usr/bin/seq
```

```
$ man seq
```

```
$ seq 3
1
2
3
```

```
$ seq 10 5 20
10
15
20
```

servewd

Serve the current working directory using a simple HTTP server. **servewd** (version 0.1) by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type servewd  
servewd is /usr/bin/dsutils/servewd  
  
$ servewd --help  
  
$ cd /data && servewd 8000
```

shuf

Generate random permutations. **shuf** (version 8.30) by Paul Eggert (2019).
More information: <https://www.gnu.org/software/coreutils>.

```
$ type shuf  
shuf is /usr/bin/shuf  
  
$ man shuf  
  
$ echo {a..z} | tr '\n' '' | shuf | trim 5  
v  
g  
m  
z  
k  
... with 21 more lines  
  
$ shuf -i 1-100 | trim 5  
13  
96  
68  
50  
46  
... with 95 more lines
```

skll

SciKit-Learn Laboratory. **skll** (version 2.5.0) by Educational Testing Service (2021). The actual tool is `run_experiment`. I use the alias `skll` because I find that easier to remember. More information:
<https://skll.readthedocs.org>.

```
$ type skill  
skill is an alias for run_experiment  
  
$ skill --help
```

sort

Sort lines of text files. `sort` (version 8.30) by Mike Haertel and Paul Eggert (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type sort  
sort is /usr/bin/sort  
  
$ man sort  
  
$ echo '3\n7\n1\n3' | sort  
1  
3  
3  
7
```

split

Split a file into pieces. `split` (version 8.30) by Torbjorn Granlund and Richard M. Stallman (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type split  
split is /usr/bin/split  
  
$ man split
```

sponge

Soak up standard input and write to a file. `sponge` (version 0.65) by Colin Watson and Tollef Fog Heen (2021). Useful if you want to read from and write to the same file in a single pipeline. More information: <https://joeyh.name/code/moreutils>.

```
$ type sponge
sponge is /usr/bin/sponge
```

sql2csv

Execute an SQL query on a database and output the result to a CSV file. **sql2csv** (version 1.0.5) by Christopher Groskopf (2020). More information: <https://csvkit.rtfd.org>.

```
$ type sql2csv
sql2csv is /usr/bin/sql2csv

$ sql2csv --help
```

ssh

OpenSSH remote login client. **ssh** (version 1:8.2p1-4ubuntu0.2) by Tatu Ylonen, Aaron Campbell, Bob Beck, Markus Friedl, Niels Provos, Theo de Raadt, and Dug Song (2020). More information: <https://www.openssh.com>.

```
$ type ssh
ssh is /usr/bin/ssh

$ man ssh
```

sudo

Execute a command as another user. **sudo** (version 1.8.31) by Todd C. Miller (2019). More information: <https://www.sudo.ws>.

```
$ type sudo
sudo is /usr/bin/sudo

$ man sudo
```

tail

Output the last part of files. `tail` (version 8.30) by Paul Rubin, David MacKenzie, Ian Lance Taylor, and Jim Meyering (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type tail  
tail is /usr/bin/tail  
  
$ man tail
```

tapkee

An efficient dimension reduction library. `tapkee` (version 1.2) by Sergey Lisitsyn, Christian Widmer, and Fernando J. Iglesias Garcia (2013). More information: <http://tapkee.lisitsyn.me>.

```
$ type tapkee  
tapkee is /usr/bin/tapkee  
  
$ tapkee --help
```

tar

An archiving utility. `tar` (version 1.30) by John Gilmore and Jay Fenlason (2014). More information: <https://www.gnu.org/software/tar>.

```
$ type tar  
tar is /usr/bin/tar  
  
$ man tar
```

tee

Read from standard input and write to standard output and files. `tee` (version 8.30) by Mike Parker, Richard M. Stallman, and David MacKenzie (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type tee  
tee is /usr/bin/tee  
  
$ man tee
```

telnet

User interface to the TELNET protocol. `telnet` (version 0.17) by Mats Erik Andersson, Andreas Henriksson, and Christoph Biedl (1999). More information: <http://www.hcs.harvard.edu/~dholland/computers/netkit.html>.

```
$ type telnet  
telnet is /usr/bin/telnet
```

tldr

Collaborative cheat sheets for console commands. `tldr` (version 3.3.7) by Owen Voke (2021). More information: <https://tldr.sh>.

```
$ type tldr  
tldr is /usr/local/bin/tldr
```

```
$ tldr --help
```

```
$ tldr tar | trim  
✓ Page not found. Updating cache...  
✓ Creating index...
```

```
tar
```

Archiving utility.

Often combined with a compression method, such as gzip or bzip2.
More information: <https://www.gnu.org/software/tar>.

- [c]reate an archive and write it to a [f]ile:
`tar cf target.tar file1 file2 file3`

... with 22 more lines

tr

Translate or delete characters. `tr` (version 8.30) by Jim Meyering (2018). More information: <https://www.gnu.org/software/coreutils>.

```
$ type tr  
tr is /usr/bin/tr  
  
$ man tr
```

tree

List contents of directories in a tree-like format. `tree` (version 1.8.0) by Steve Baker (2018). More information: <https://launchpad.net/ubuntu/+source/tree>.

```
$ type tree  
tree is /usr/bin/tree  
  
$ man tree  
  
$ tree -L 1 | trim  
/  
├── bin -> usr/bin  
├── boot  
└── data  
    ├── ch01  
    ├── ch02  
    │   └── fac.py  
    │   └── movies.txt  
    └── ch03  
        └── logs.tar.gz  
... with 122908 more lines
```

trim

Trim output to a given height and width. `trim` by Jeroen Janssens (2021). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type trim  
trim is /usr/bin/dsutils/trim  
  
$ echo {a..z}{0..9} | fold | trim 5 60  
a-0 a-1 a-2 a-3 a-4 a-5 a-6 a-7 a-8 a-9 b-0 b-1 b-2 b-3 b-4...  
c-0 c-1 c-2 c-3 c-4 c-5 c-6 c-7 c-8 c-9 d-0 d-1 d-2 d-3 d-4...  
e-0 e-1 e-2 e-3 e-4 e-5 e-6 e-7 e-8 e-9 f-0 f-1 f-2 f-3 f-4...  
g-0 g-1 g-2 g-3 g-4 g-5 g-6 g-7 g-8 g-9 h-0 h-1 h-2 h-3 h-4...  
i-0 i-1 i-2 i-3 i-4 i-5 i-6 i-7 i-8 i-9 j-0 j-1 j-2 j-3 j-4...  
... with 8 more lines
```

ts

Timestamp input. **ts** (version 0.65) by Joey Hess (2021). More information: <https://joeyh.name/code/moreutils>.

```
$ type ts  
ts is /usr/bin/ts  
  
$ echo seq 5 | sample -d 500 | ts  
Jun 29 14:39:19 seq 5
```

type

Show the type and location of a command-line tool. **type** is a Z shell builtin.

```
$ type type  
type is a shell builtin  
  
$ man zshbuiltins | grep -A 10 '^ *type '
```

uniq

Report or omit repeated lines. **uniq** (version 8.30) by Richard M. Stallman and David MacKenzie (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type uniq  
uniq is /usr/bin/uniq
```

```
$ man uniq
```

unpack

Extract common file formats. **unpack** (version 0.1) by Patrick Brisbin (2013). More information: <https://github.com/jeroenjanssens/dsutils>.

```
$ type unpack  
unpack is /usr/bin/dsutils/unpack
```

unrar

Extract files from RAR archives. **unrar** (version 0.0.1) by Ben Asselstine, Christian Scheurer, and Johannes Winkelmann (2014). More information: <https://web.archive.org/web/20080331080828/http://home.gna.org/unrar/>.

```
$ type unrar  
unrar is /usr/bin/unrar
```

```
$ man unrar
```

unzip

List, test, and extract compressed files in a ZIP archive. **unzip** (version 6.0) by Samuel H. Smith, Ed Gordon, Christian Spieler, Onno van der Linden, Mike White, Kai Uwe Rommel, Steven M. Schweda, Paul Kienitz, Chris Herborth, Jonathan Hudson, Sergio Monesi, Harald Denker, John Bush, Hunter Goatley, Steve Salisbury, Steve Miller, and Dave Smith (2009). More information: <http://www.info-zip.org/pub/infozip>.

```
$ type unzip  
unzip is /usr/bin/unzip
```

```
$ man unzip
```

VW

Fast machine learning library for online learning. **vw** (version 8.10.1) by John Langford (2021). More information: <https://vowpalwabbit.org>.

```
$ type vw
vw is /usr/local/bin/vw

$ vw --help --quiet
```

WC

Print newline, word, and byte counts for each file. **wc** (version 8.30) by Paul Rubin and David MacKenzie (2019). More information: <https://www.gnu.org/software/coreutils>.

```
$ type wc
wc is /usr/bin/wc

$ man wc
```

which

Locate a command. **which** (version 0.1) by unknown (2016).

```
$ type which
which is a shell builtin

$ man which
```

xml2json

Convert an XML input to a JSON output, using xml-mapping. **xml2json** (version 0.0.3) by François Parmentier (2016). More information: <https://github.com/parmentf/xml2json>.

```
$ type xml2json  
xml2json is /usr/local/bin/xml2json
```

xmlstarlet

Command-line XML/XSLT toolkit. `xmlstarlet` (version 1.6.1) by Dagobert Michelsen, Noam Postavsky, and Mikhail Grushinskiy (2019). More information: <https://sourceforge.net/projects/xmlstar>.

```
$ type xmlstarlet  
xmlstarlet is /usr/bin/xmlstarlet  
  
$ man xmlstarlet
```

xsv

A fast CSV command-line toolkit written in Rust. `xsv` (version 0.13.0) by Andrew Gallant (2018). More information: <https://github.com/BurntSushi/xsv>.

```
$ type xsv  
xsv is /usr/bin/xsv  
  
$ xsv --help
```

zcat

Decompress and concatenate files to standard output. `zcat` (version 1.10) by Paul Eggert (2021). More information: <https://www.nongnu.org/zutils/zutils.html>.

```
$ type zcat  
zcat is /usr/bin/zcat  
  
$ man zcat
```

zsh

The Z shell. **zsh** (version 5.8) by Paul Falstad and Peter Stephenson (2020).
More information: <https://www.zsh.org>.

```
$ type zsh
zsh is /usr/bin/zsh
```

```
$ man zsh
```

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About the Author

Jeroen Janssens is an independent data science consultant and instructor. He enjoys visualizing data, implementing machine learning models, and building solutions using Python, R, JavaScript, and Bash. Jeroen manages **Data Science Workshops**, a training and coaching firm that organizes open enrollment workshops, in-company courses, inspiration sessions, hackathons, and meetups. Previously, he was an assistant professor at Jheronimus Academy of Data Science and a data scientist at Elsevier in Amsterdam and various startups in New York City. Jeroen holds a PhD in machine learning from Tilburg University and an MSc in artificial intelligence from Maastricht University. He lives with his wife and two kids in Rotterdam, the Netherlands.

Colophon

The animal on the cover of *Data Science at the Command Line* is a wreathed hornbill (*Rhytidoceros undulatus*). Also known as the bar-pouched wreathed hornbill, the species is found in forests in mainland Southeast Asia and in northeastern India and Bhutan. Hornbills are named for the *casques* that form on the upper part of the birds' bills. No single obvious purpose exists for these hollow, keratinized structures, but they may serve as a means of recognition between members of the species, as an amplifier for the birds' calls, or—because males often exhibit larger casques than females of the species—for gender recognition. Wreathed hornbills can be distinguished from plain-pouched hornbills, to whom they are closely related and otherwise similar in appearance, by a dark bar on the lower part of the wreathed hornbills' throats.

Wreathed hornbills roost in flocks of up to four hundred but mate in monogamous, lifelong partnerships. With help from the males, females seal themselves up in tree cavities behind dung and mud to lay eggs and brood. Through a slit large enough for his beak alone, the male feeds his mate and their young for up to four months. A diet of animal prey becomes predominantly fruit when females and their young leave the nest. Hornbill couples have been known to return to the same nest for as many as nine years.

Many of the animals on O'Reilly covers are endangered; all of them are important to the world.

The color illustration is by Karen Montgomery, based on a black and white engraving from Braukhaus's *Lexicon*. The cover fonts are Gilroy Semibold and Guardian Sans. The text font is Adobe Minion Pro; the heading font is Adobe Myriad Condensed; and the code font is Dalton Maag's Ubuntu Mono.

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