

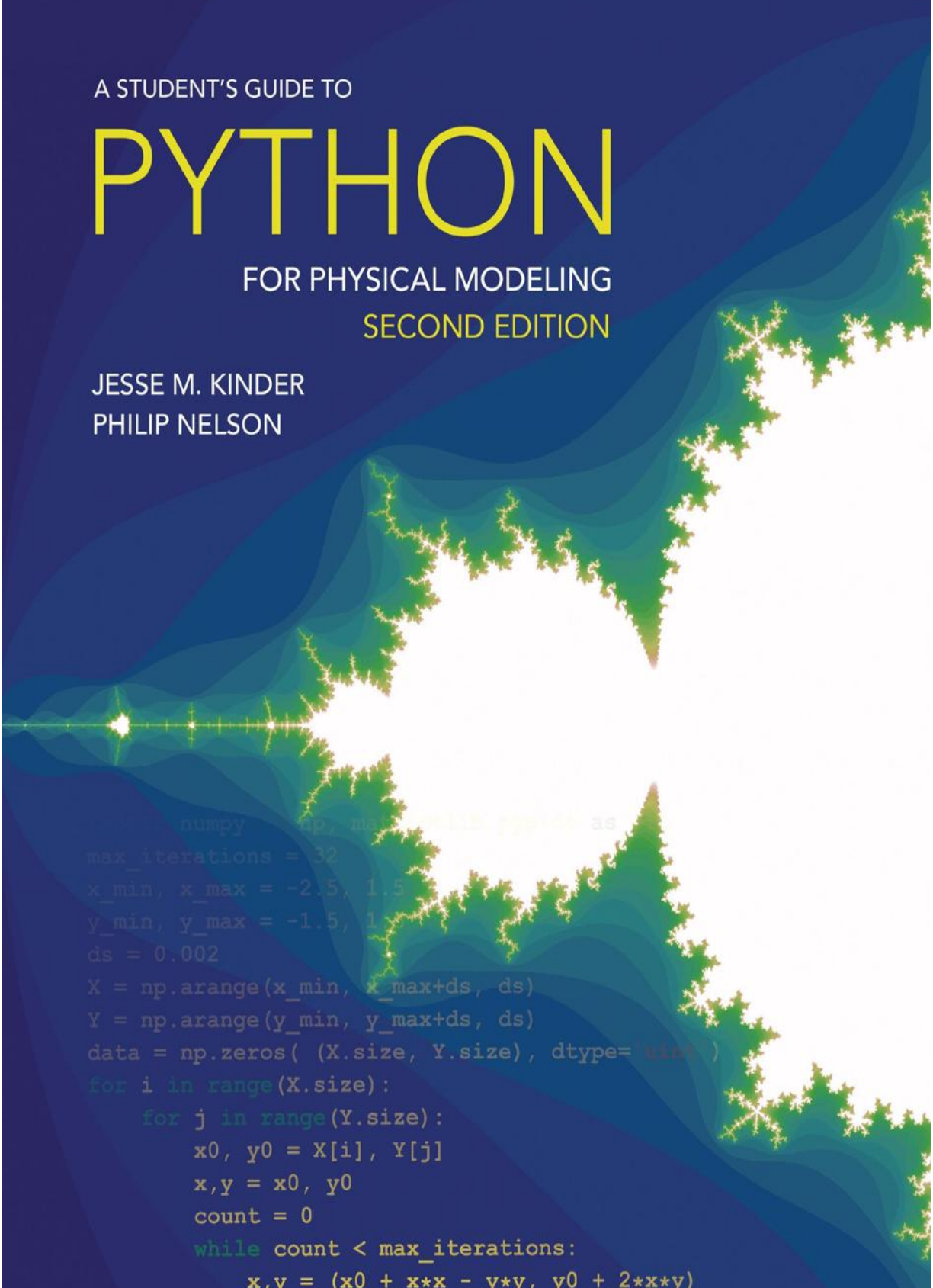
A STUDENT'S GUIDE TO

PYTHON

FOR PHYSICAL MODELING

SECOND EDITION

JESSE M. KINDER
PHILIP NELSON



```
import numpy as np, math, pylab as pl
max_iterations = 32
x_min, x_max = -2.5, 1.5
y_min, y_max = -1.5, 1
ds = 0.002
X = np.arange(x_min, x_max+ds, ds)
Y = np.arange(y_min, y_max+ds, ds)
data = np.zeros( (X.size, Y.size), dtype= uint8 )
for i in range(X.size):
    for j in range(Y.size):
        x0, y0 = X[i], Y[j]
        x,y = x0, y0
        count = 0
        while count < max_iterations:
            x,y = (x0 + x*x - y*y, y0 + 2*x*y)
            if abs(x) > 2 or abs(y) > 2:
                break
            count += 1
        data[i,j] = count
pl.imshow(data, origin='lower')
pl.show()
```


A Student's Guide to Python for Physical Modeling

Second Edition

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Jesse M. Kinder and Philip Nelson

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Princeton and Oxford

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versions of Python or its extended environments.

For Oliver Arthur Nelson—PN

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Let's Go

Why teach yourself Python, and why do it this way

Learning to program a computer can change your perspective. At first, it feels like you are struggling along and picking up a couple of neat tricks here and there, but after a while, you start to realize that *you* can make the computer do almost *anything*. You can add in the effects of friction and air resistance that your physics professor is always telling you to ignore, you can make your own predator-prey simulations to study population models, you can create your own fractals, you can look for correlations in the stock market—the list is endless.

In order to communicate with a computer, you must first learn a language it understands. Python is an excellent choice, because it is easy to get started and its structure is very natural—at least compared to some other computer languages. Soon, you will find yourself spending most of your time thinking about how to solve a problem, rather than how to explain your calculation to a computer.

Whatever your motivation for learning Python, you may wonder whether it's really necessary to wade through everything in this book. Bear with us. We are working scientists, and we have used our experience to prepare you to start exploring and learning on your own as efficiently as possible. Spend a few hours trying everything we recommend, in the order we recommend it. This will save time in the long run. We have eliminated everything you don't need at the outset. What remains is a set of basic knowledge and skills that you will almost certainly find useful someday.

How to use this tutorial

Here are a few ideas about how you might teach yourself Python using this book.

- Many code samples that appear in this document, as well as errata, updates, data sets, and more are available via press.princeton.edu/titles/32489.html. Use these resources.
- After the first few pages, you'll need to work in front of a computer that is running Python. (Don't worry—we'll tell you how to set it up.) On that same computer, you'll probably want to have open a text document named `code_samples.txt`, which is available via the website above.
- Next to the computer you may have a hard copy of this book, or the eBook on a tablet or other device. Alternatively, the eBook can be open on the same computer that runs Python.
- This book will frequently ask you to try things. Some of those things involve snippets of code given in the text. You can copy and paste code from `code_samples.txt` into your Python session, see what happens, then modify and play with it.
- You can also access the snippets interactively. A page with links to individual code samples is available via the website above. In the eBook, you can also click on the words [get code] at the top of a code sample to visit the web page. Either way, you can copy and paste code from the web page into Python.
- A few sections and footnotes are flagged with this “Track 2” symbol: . These are more advanced and can be skipped on a first reading.

And now ... Let's go.

A Student's Guide to Python for Physical Modeling

Second Edition

CHAPTER 1

Getting Started with Python

*The Analytical Engine weaves algebraical patterns,
just as the Jacquard loom weaves flowers and leaves.*

— Ada, Countess of Lovelace, 1815–1853

1.1 ALGORITHMS AND ALGORITHMIC THINKING

The goal of this tutorial is to get you started in computational science using the computer language Python. Python is open-source software. You can download, install, and use it anywhere. Many good introductions exist, and more are written every year. *This one* is distinguished mainly by the fact that it focuses on skills useful for solving problems in physical modeling.

Modeling a physical system can be a complicated task. Let's take a look at how we can use the powerful processors inside your computer to help.

1.1.1 Algorithmic thinking

Suppose that you need to instruct a friend how to back your car out of your driveway. Your friend has never driven a car, but it's an emergency, and your only communication channel is a phone conversation before the operation begins.

You need to break the required task down into small, explicit steps that your friend understands and can execute in sequence. For example, you might provide your friend with the following set of instructions:

- 1 Put the key in the ignition.
- 2 Turn the key until the car starts, then let go.
- 3 Push the button on the shift lever and move it to "Reverse."
- 4 ...

Unfortunately, for many cars this “code” won’t work, even if your friend understands each instruction: It contains a **bug**. Before step 3, many cars require that the driver

Press down the left pedal.

Also, the shifter may be marked R instead of Reverse. It is difficult at first to get used to the high degree of precision required when composing instructions like these.

Because you are giving the instructions in advance (your friend has no mobile phone), it’s also wise to allow for contingencies:

If a crunching sound is heard, press down on the left pedal ...

2 Chapter 1 Getting Started with Python

Breaking the steps of a long operation down into small, explicit substeps and anticipating contingencies are the beginning of *algorithmic thinking*.

If your friend has had a lot of experience watching people drive cars, then the instructions above may suffice. But a friend from Mars—or a robot—would need much more detail. For example, the first two steps may need to be expanded to something like

```
Grab the wide end of the key.  
Insert the pointed end of the key into the slot on the lower right side  
of the steering column.  
Rotate the key about its long axis in the clockwise direction  
(when viewed from the wide end toward the pointed end).  
...
```

These two sets of instructions illustrate the difference between low-level and high-level languages for communicating with a computer. A *low-level* computer program is similar to the second set of explicit instructions, written in a language that a machine can understand.¹ A *high-level* system understands many common tasks, and therefore can be programmed in a more condensed style, like the first set of instructions above. Python is a high-level language. It includes commands for common operations in mathematical calculations, processing text, and manipulating files. In addition, Python can access many *standard libraries*, which are collections of programs that perform advanced functions such as data visualization and image processing.

Python also comes with a *command line interpreter*—a program that executes Python commands as you type them. Thus, with Python, you can save instructions in a file and run them later, or you can type commands and execute them immediately. In contrast, many other programming languages used in scientific computing, like C++ or FORTRAN, require you to *compile* your programs before you can *execute* them. A separate program called a compiler translates your code into a low-level language. You then run the resulting compiled program to execute (carry out) your algorithm. With Python, it is comparatively easy to quickly write, run, and debug programs. (It still takes patience and practice, though.)

A command line interpreter combined with standard libraries and programs you write yourself provides a convenient and powerful scientific computing platform.

1.1.2 States

You have probably studied multistep mathematical proofs, perhaps long ago in a geometry class. The goal is to verify a proposition through a chain of steps that use given information and a formal system. Thus, each statement's truth, although not evident in isolation, is supposed to be straightforward in light of the preceding statements. The reader's “state” (list of propositions known to be true) changes while reading through the proof. At the end, that list includes the desired result.

An **algorithm** has a different goal. It is a chain of instructions, each of which describes a simple operation, that accomplishes a complex task. The chain may involve a lot of repetition, so you won't want to supervise the execution of every step. Instead, you specify all the steps in advance, then stand back while your electronic assistant performs them rapidly. There may also be contingencies that cannot be known in advance. (*If a crunching sound is heard, ...*)

In an algorithm, the *computer* has a state that is constantly being modified. For example, it has many memory cells, whose contents may change during the course of an operation. Your goal might be to

¹ Machine code and assembly language are low-level programming languages.

arrange for one or more of these cells to contain the result of some complex calculation once the algorithm has finished running. You may also want a particular graphical image to appear.

1.1.3 What does `a = a + 1` mean?

To get a computer to execute your algorithm, you must first express it in a programming language. The commands used in computer programming can be confusing at first, especially when they contradict standard mathematical usage. For example, many programming languages (including Python) accept statements such as these:

```
1 a = 100
2 a = a + 1
```

In mathematics, this makes no sense. The second line is an assertion that is always false; equivalently, it is an equation with no solution. To Python, however, “=” is not a test of equality, but an instruction to be executed. These lines have roughly the following meaning:²

1. Assign the name `a` (a **variable**) to an integer object with the value 100.
2. Extract the value of the object named `a`. Calculate the sum of that value and 1. Assign the name `a` to the result, and *discard* whatever was previously stored under the name `a`.

In other words, the equals sign instructs Python to change its *state*. In contrast, mathematical notation uses the equals sign to create a proposition, which may be true or false. Note, too, that Python treats the left and right sides of the command `x=y` differently, whereas in math the equals sign is symmetric. For example, Python will give an error message if you say something like `b+1=a`; the left side of an assignment must be a name that can be assigned to the result of evaluating the right side.

We do often wish to check whether a variable has a particular value. To avoid ambiguity between assignment and testing for equality, Python uses a double equals sign for the latter:

```
1 a = 1
2 a == 0
3 b = (a == 1)
```

This code again creates a variable `a` and assigns it to a numerical value. Then it compares this numerical value with 0. Finally, it creates a second variable `b`, and assigns it a logical value (**True** or **False**) after performing another comparison. That value can be used in contingent code, as we'll see later.

Do not use `=` (assignment) when `==` (test for equality) is required.

This is a common mistake for beginning programmers. You can get mysterious results if you make this error, because both `=` and `==` are legitimate Python syntax. In any particular situation, however, only one of them is what you want.

1.1.4 Symbolic versus numerical

In math, it's perfectly reasonable to start a derivation with “Let $b = a^2 - a$,” even if the reader doesn't yet know the value of a . This statement defines b in terms of a , whatever the value of a may be.

² [T2](#) Appendix F gives more precise information about the handling of assignment statements.

4 Chapter 1 Getting Started with Python

If you launch Python and immediately give the equivalent statement, `b=a**2-a`, the result is an error message.³ Every time you hit <Return/Enter>, Python tries to compute values for every assignment statement. If the variable `a` has not been assigned a value yet, evaluation fails, and Python complains. Other computer math packages can accept such input, keep track of the symbolic relationship, and evaluate it later, but basic Python does not.⁴

In math, it's also understood that a definition like “Let $b = a^2 - a$ ” will persist unchanged throughout the discussion. If we say, “In the case $a = 1, \dots$ ” then the reader knows that b equals zero; if later we say, “In the case $a = 2, \dots$ ” then we need not reiterate the definition of b for the reader to know that this symbol now represents the value $2^2 - 2 = 2$.

In contrast, a numerical system like Python *forgets* any relation between `b` and `a` after executing the assignment `b=a**2-a`. All that it remembers is the *value* now assigned to `b`. If we later change the value of `a`, the value of `b` will *not* change.⁵

Changing symbolic relationships in the middle of a proof is generally not a good idea. However, in Python, if we say `b=a**2-a`, nothing stops us from later saying `b=2**a`. The second assignment updates Python's state by discarding the value calculated in the first assignment statement and replacing it with the newly computed value.

1.2 LAUNCH PYTHON

Rather than reading about what happens when you type some command, try out the commands for yourself. Appendix A describes how to install and launch Python. From now on, you should have Python running as you read: Try every snippet of code and observe what Python does in response. For example, this tutorial won't show you much graphics or output. You must generate these yourself as you work through the examples.

*Reading this tutorial won't teach you Python. You can **teach yourself** Python by working through all the examples and exercises here, and then using what you've learned on your own problems.*

Set yourself little challenges and test them out. (“What would happen if …?” “How could I accomplish…?”) Python is not some expensive piece of lab apparatus that could break or explode if you type something wrong! Just try things. This strategy is not only more fun than passively accumulating facts—it is also far more effective.

Before you start typing, we would like to explain a few conventions we use in this book. The most important is this:

Python code consists entirely of plain text.

All fonts, typefaces, and coloring in the code samples of this tutorial were added for readability. These are not things you need to worry about while entering code. Similarly, the line numbers shown on the left of code samples are there to allow us to refer to particular lines. Don't type them. Spyder will assign and

³ The notation `**` denotes exponentiation. See Section 1.4.2.

⁴ The SymPy library makes symbolic calculations possible in Python. See Section 10.3.2.

⁵ In math, the statement $b = a^2 - a$ essentially defines b as a *function* of a . We can certainly do that in Python by defining a function

that returns the value of $a^2 - a$ and assigning that function the name `b` (see Section 6.1), but this is *not* what “=” does.

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show line numbers when you work in the Editor, and Python will use them to tell you where it thinks you have made errors. They are not part of the code. Note also that most blank spaces are optional, except when used for indentation. We use extra blank spaces to improve readability, but these are not required.

This tutorial uses the following color and font scheme when displaying code:

- Built-in functions and reserved words are displayed in boldface green type:
`print("Hello, world!")`. You do not need to import these functions.
- Python errors and runtime exceptions are displayed in boldface red type: `SyntaxError`.
- Functions and other objects from NumPy and PyPlot are displayed in boldface black type: `np.sqrt(2)`, or `plt.plot(x,y)`. We will assume that you import NumPy and PyPlot at the beginning of each session and program you write.⁶
- Functions imported from other libraries are displayed in blue boldface type:
`from scipy.special import factorial`.
- Strings are displayed in red type: `print("Hello, world!")`.
- Comments are displayed in oblique blue type: `# This is a comment`.
- Keywords in function arguments are displayed in oblique black type:
`np.loadtxt('data.csv', delimiter=',')`. Keywords are not arbitrary; they must be spelled exactly as shown.
- Buttons you can click with a mouse are displayed in small capitals within a rectangle: . Some buttons in Spyder have icons rather than text, but hovering the mouse pointer over the button will display the text shown in this tutorial.
- Keystrokes are displayed within angled brackets: `<Return>` or `<Ctrl-C>`.
- Most other text is displayed in normal type.

Regarding keystrokes, our notation may not look exactly like what you see on your keyboard, so we summarize our conventions in the table below. All keys that appear in a single unit should be pressed together. For example, `<Ctrl-C>` means press and hold the “control” key on your keyboard, and while holding it press the “c” key (you may also see this abbreviated as `^C`). Our conventions follow the macOS keyboard layout. If you are using Windows or Linux, substitute `<Ctrl>` for `<Cmd>`. We’ll abbreviate `<Return/Enter>` as simply `<Return>`.

Key	Example	Function
enter or return	<code><Return></code>	end line or run command
control	<code><Ctrl-C></code>	interrupt current Python command
command	<code><Cmd-V></code>	paste from the clipboard
option or alt	<code><Alt-Shift-R></code>	restart Spyder

Now that you know what to type (plain text) and how to type it, all you need is Python! A complete Python programming environment has many components. See Table 1.1 for a brief description of the ones that we’ll be discussing. Be aware that we use “Python” loosely in this guide. In addition to the language itself, Python may refer to a *Python interpreter*, which is a computer application that accepts commands and performs the steps described in a program. Python may also refer to the language together with common libraries.

⁶ See Section 1.3 (page 11).

6 Chapter 1 Getting Started with Python

Python	A computer programming language. A way to describe algorithms to a computer.
IPython	A Python <i>interpreter</i> : A computer application that provides a convenient, interactive mode for executing Python commands and programs.
Spyder	An <i>integrated development environment</i> (IDE): A computer application that includes IPython, a tool to inspect variables, a text editor for writing and debugging programs, and more.
Jupyter	A notebook-style interface for Python.
NumPy	A standard library that provides numerical arrays and mathematical functions.
PyPlot	A standard library that provides visualization tools.
SciPy	A standard library that provides scientific computing tools.
Anaconda	A <i>distribution</i> : A single download that includes all of the above and provides access to many additional libraries for special purposes. It also includes a <i>package manager</i> that helps you to keep everything up to date.

Table 1.1: Elements of the Python environment described in this tutorial.

Most of the code that follows will run with any Python distribution. However, since we cannot provide instructions for every available version of Python and every integrated development environment (IDE), we have chosen the following particular setup:

- The Anaconda distribution of Python 3, available at anaconda.com. Many scientists instead use an earlier version of Python (such as version 2.7). Appendix E discusses the minor changes needed to adapt the codes in this tutorial for earlier versions.
- The Spyder IDE, which comes with Anaconda or can be obtained at www.spyder-ide.org. Any programming task can be accomplished with a different IDE—or with no IDE at all. Other IDEs are available, such as IDLE, which comes with every distribution of Python. Browser-based Jupyter notebooks and JupyterLab are another option.⁷

The choice of distribution is a matter of personal preference. We chose Anaconda because it is simple to install, update, and maintain, and it is free. You may find a different distribution is better suited to your needs. For example, you can install Python from source (python.org) and manage your packages with pip, but this tutorial will assume you are using Anaconda and the conda package manager.

1.2.1 IPython console

To keep our discussion focused on Python rather than on details of various platforms, we will assume that you are using Spyder as you work through this guide. This is not required! If you prefer to start with a simpler interface, you can open the Qt Console app from the Anaconda Navigator and start typing commands. If you prefer a notebook interface, you can follow along in a Jupyter Notebook. (See Appendix C.) If you prefer working from the command line, you can start IPython from a terminal. (See Appendix B.) At some point, though, you will need an IPython interpreter and a text editor. Spyder includes both of these, plus some other useful features, in an interface that will be familiar to users of MATLAB. There are many ways to use Python, and you can use any of them as you work through this tutorial. If you are new to Python, Spyder is a good choice.

Open Spyder now. Upon launch, Spyder opens a window that includes several *panes*. See Figure 1.1.

There is an Editor pane on the left for editing program files (*scripts*). There are two panes on the right.

⁷ If you prefer the notebook interface, see Appendix C to get started. Many code samples are available in notebook format via this book's blog.

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1.2 Launch Python 7

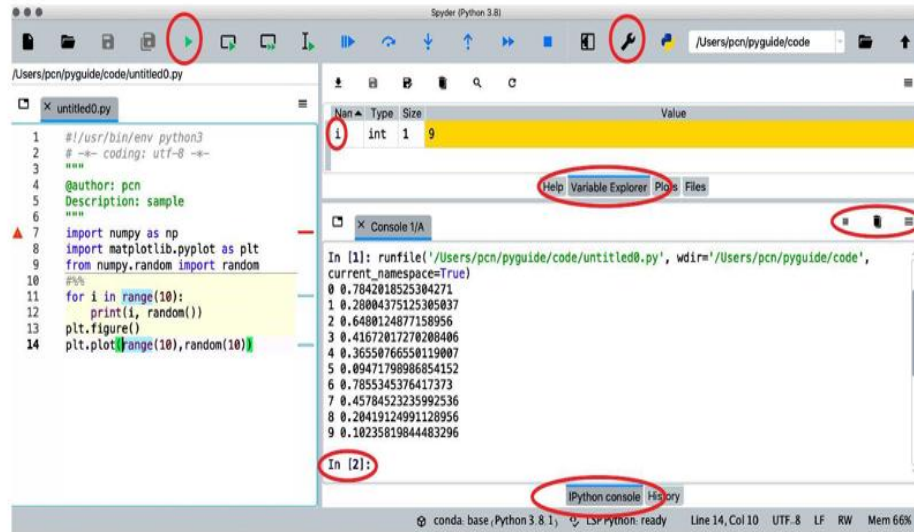


Figure 1.1: The Spyder display. Red circles have been added to emphasize (from top to bottom) the **RUN** button, Preferences (wrench icon), a variable in the Variable Explorer, the tab that brings the Variable Explorer to the front in its pane, the **STOP**, **RESET**, and **OPTIONS** buttons, the IPython command prompt, and the IPython console tab. Two scripts are open in the Editor; *untitled0.py* has been brought to the front by clicking its tab at the top of the Editor pane. Two warnings appear at far left.

The top-right pane may contain Help, Variable Explorer, Plots, and File Explorer tabs. If necessary, click on the Variable Explorer's tab to bring it to the front. The bottom-right pane should include a tab called "IPython console"; if necessary, click it now.⁸ It provides the command line interpreter that allows you to execute Python commands interactively as you type them.

If your window layout gets disorganized, do not worry. It is easy to adjust. The standard format for Spyder is a single window, divided into the three panes just described. Each pane can have multiple tabs. If you have unwanted windows, close them individually by clicking on their CLOSE buttons. You can also use the menu **View>Panes** to select panes you want to be visible and deactivate those you do not want. **View>Window layouts>Spyder Default Layout** will restore the standard layout.

Click in the IPython console. Now, things you type will show up after the *command prompt*. By default, this will be something like

In[1]:

Try typing simple commands like "2+2" and hitting <Return> after each line. Python responds immediately after each <Return>, attempting to perform whatever command you entered.⁹

Click on the Variable Explorer tab. Each time you enter a command and hit <Return>, the contents

⁸ If no IPython console tab is present, you can open one from the menu at the top of the screen:
Consoles>New console.

⁹ This tutorial uses the word “command” to mean any Python statement that can be executed by an interpreter. Assignments like `a=1`, function calls like `plt.plot(x,y)`, and special instructions like `%reset` are commands.

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8 Chapter 1 Getting Started with Python

of this pane will reflect any changes in Python’s state: Initially empty, it will display a list of your variables and a summary of their values.¹⁰ When a variable contains many values (for example, an array), you can double-click its entry in this list to open a spreadsheet that contains all the values of the array. You can copy from this spreadsheet and paste into other applications.

At any time, you can reset Python’s state by quitting and relaunching it, or by executing the command

`%reset`

Because you are about to delete almost everything that has been created in this session, you will be asked to confirm this irreversible operation.¹¹ Press `<y>` then `<Return>` to proceed. (Commands that begin with a `%` symbol are **magic commands**: commands specific to the IPython interpreter. They may not work in a more basic Python interpreter, or in scripts that you write. To learn more about these, type `%magic` at the IPython command prompt.)

Example: Use the `%reset` command, then try the following commands at the prompt. Type each line exactly as shown, then press `<Return>`. Explain everything you see happen:

```
q  
q == 2  
q = 2  
q  
q == 2  
q == 3
```

Solution: Python complains about the first two lines: Initially, the symbol `q` is not associated with any object. It has no value, and so expressions involving it cannot be evaluated. Altering Python’s state in the third line above changes this situation, so the last three lines do not generate errors.

Example: Now clear Python’s state again. Try the following at the prompt, and explain everything that happens. (It may be useful to refer to Section 1.1.4.)

```
a = 1  
a  
b = a**2 - a  
b  
a = 2  
print(a)  
print(b)  
b = a**2 - a  
a, b  
print(a, b)
```

Solution: The results from the first four lines should be clear: We assign values to the variables `a` and `b`.

In the fifth line, we change the value of `a`, but because Python remembers only the *value* of `b` and not its relation to `a`, `b`'s value is unchanged until we update it explicitly in the eighth line.

¹⁰ Some variables will not appear. You can control which variables are excluded through the `OPTIONS` menu, in the upper-right corner of the Variable Explorer pane.

¹¹ If IPython does not seem to respond to `%reset`, try scrolling the IPython console up manually to see the confirm query.

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1.2 Launch Python 9

When entering code at the command prompt, you may run into a confusing situation where Python seems unresponsive and displays “`...:`” instead of executing commands.

If a command contains an unmatched (, [, or {, then Python continues reading more lines, searching for the corresponding),], or }.

Look for an unmatched bracket. If you find one, type the closing bracket and press `<Return>`. If you cannot figure out how to match up your brackets, or if there is some other problem, you can force execution with `<Shift + Return>` or abort the command by pressing `<Esc>`.¹²

The examples above illustrate an important point: An assignment statement does not display the value that it assigns to a variable. To see the value assigned to a variable in an IPython session, use the `print()` command or type the variable name on a line by itself.¹³

The last two lines of the example above illustrate how to see the values of multiple objects at once. Notice that the output is not exactly the same.

You can end a command by starting a new line. Or, if you wish, you can end a command with a semicolon (`;`) and then add another command on the same line.

It is also possible to make multiple assignments with a single `=` command. This is an alternative to using semicolons. Both of the following lines assign the same values to their respective variables:

```
a = 1; b = 2; c = 3
x, y, z = 1, 2, 3
```

Either side of the second command may be enclosed in parentheses without affecting the result.

The preceding paragraph demonstrates ways to save space and reduce typing with Python. Sometimes this is convenient, but it's best not to make too much use of this ability. You should instead try to make the *meaning* of your code as clear as possible. Human readability is worth a few extra seconds of typing or a few extra lines in a program.

In some situations, you may wish to use a very long command that doesn't fit on one line. For such cases, you can end a line with a backslash (`\`). Python will then continue reading the next line as part of the same command. Try this:

```
q = 1 + \
2
print(q)
```

A single command can even stretch over multiple lines:

```
xv\
a\
1\
= 1 + \
2
```

This will create a variable `xval` and assign it the value 3.

To write clear code, use backslashes and semicolons sparingly.

¹² `<Esc>` cancels the current command in Spyder. In another IDE or interpreter, you may need to use `<Ctrl-C>` to interrupt and `<Alt+Return>` to force execution.

¹³ In scripts that you write, Python will evaluate an expression *without* showing anything on the screen; if you want output, you must give an explicit `print()` command. Scripts will be discussed in Section 3.3.

1.2.2 Error messages

You should have encountered some error messages by now. When Python detects an error, it tells you where it encountered the error, provides a fragment of the code surrounding the statement that caused the problem, and tells you which general kind of error it detected among the many types it recognizes. For example, Python responds with a **NameError** whenever you try to evaluate an undefined variable. (Recall the Example on page 8.) See Appendix D for a description of common Python errors and some hints for interpreting the resulting messages.

Donald Knuth, a well-known computer scientist, once wrote, “Error messages can be terrifying when you aren’t prepared for them; but they can be fun when you have the right attitude. Just remember that you really haven’t hurt the computer’s feelings, and that nobody will hold the errors against you.” We encourage you to adopt this attitude.

- Don’t be afraid to make errors. It’s very hard to break anything.
- Read the error messages. They tell you what kind of error you made—not just that you made an error.
- Inspect the code that produces an error. You can learn from your mistakes.

This approach to errors will make you a better coder right away, and it will help you “debug” more complicated programs later.¹⁴

1.2.3 Sources of help

The definitive documentation on Python is available online at www.python.org/doc. However, in many cases you’ll find the answers you need more quickly by other means, such as asking a friend, searching the web, or visiting [stack overflow.com](http://stackoverflow.com).

Suppose that you wish to evaluate the square root of 2. You type `2**0.5` and hit <Return>. That does the job, but Python is displaying 16 digits after the decimal point, and you only want 3. You think there’s probably a function called `round` in Python, but you are not sure how to use it or how it works. You can get help directly from Python by typing `help(round)` at the command prompt. You’ll see that this is indeed the function you were looking for:

```
round(2**0.5, 3)
```

gives the desired result.

In Spyder, there are additional ways to get help. Type `round` at the command prompt, but do not hit <Return>. Instead hit <Cmd-I> or <Ctrl-I> (for “Information”). The information that was displayed in the IPython console when you issued the `help` command now shows up in the Help tab, and in a format that is easier to navigate and read, especially for long entries. You can also use the Help tab without entering anything at the command prompt: Try entering `pow` in the “Object” field at the top of the pane. The Help tab provides information about an alternative to `**` for raising a number to a power.

In IPython, you can also follow or precede the name of any Python object, including function and variable names, by a question mark to obtain help: `round?` or `?round` provides almost the same information as `help(round)` and is easier to type.

When you type `help(...)`, Python will print out the information it has about the expression in parentheses if it recognizes the name. Unfortunately, Python is not as friendly if you don’t know the name of the command you need. Perhaps you think there ought to be a way to take the square root of a number

¹⁴ See Section 3.3.3 (page 43).

without using the power notation. After all, it is a pretty basic operation. Type `help(sqrt)` to see what happens when Python does not recognize the name you request.

To find out what commands are currently available to you, you can use Python's `dir()` command. This is short for “`directory`,” and it returns a list of all the modules, functions, and variable names that have been created or imported during the current session (or since the last `%reset` command). Ask Python for help on `dir` to learn more. Nothing in the output of `dir()` looks like a square root, but there is an item called `__builtins__`. This is the collection of all the functions and other objects that Python recognizes when it first starts up. It is Python's “last resort” when hunting for a function or variable.¹⁵ To see the list of built-in functions, type

```
dir(__builtins__)
```

There is no `sqrt` function or anything like it. In fact, *none* of the standard mathematical functions, such as `sin`, `cos`, or `exp` show up!

Python cannot help you any further at this point. You now have to turn to outside resources. Good options include books about Python, search engines, friends who know more about Python than you do, and so on.

In the beginning, a lot of your coding time will be spent using a search engine to get help.

The `sqrt` function we seek belongs to a library. Later we will discuss how to access libraries of useful functions that are not automatically available with Python.

Your
Turn
1A

Before proceeding, try a web search for
how to take square roots in python

1.2.4 Good practice: Keep a log

As you work through this tutorial, you will hit many small roadblocks—and some large ones. How do you evaluate a modified Bessel function? What do you do if you want a subscript in a graph axis label? The list is endless. Every time you resolve such a puzzle (or a friend helps you), *make a note of how you did it* in a notebook or in a dedicated file somewhere on your computer. Later, looking through that log will be much easier than scanning through all the code you wrote months ago (and less irritating than asking your friend over and over).

1.3 PYTHON MODULES

We discovered that Python does not have a built-in `sqrt` function. Even your calculator has that! What good is Python? Think for a moment about how, exactly, your calculator knows how to find square roots. At some point in the past, someone came up with an algorithm for computing the square root of a number and stored it in the permanent memory of your calculator. Someone had to create a *program* to calculate square roots.

¹⁵ Appendix F explains how Python searches for variables and other objects.

Python is a programming language. A Python interpreter understands a basic set of commands that can be combined to perform complex tasks. Python also has a large community of developers who have created entire libraries of useful functions. To gain access to these, however, you need to **import** them into your working environment.

*Use **import** to access functions that do not come standard with Python.*

1.3.1 `import`

At the command prompt, type

```
import numpy
```

and hit <Return>. You now have access to many useful functions. You have imported the NumPy module, a collection of tools for numerical calculation using Python: “Numerical Python.” (Do not capitalize its name in your code.)

To see what has been gained, type `dir(numpy)`. You will find nearly 600 new options at your disposal, and one of them is the `sqrt` function you originally sought. You can search for the function within NumPy by using the command `numpy.lookfor('sqrt')` (This will often return more than you need, but the first few lines can be quite helpful.) Now that you have imported NumPy, try

```
sqrt(2)
```

What's going on? You just imported a square root function, but Python tells you that `sqrt` is not defined! Try instead

```
numpy.sqrt(2)
```

The `sqrt` function you want “belongs” to the `numpy` module you imported. Even after importing, you still have to tell Python where to find it before you can use it.

After you have imported a module, call its functions by giving the module name, a period, and then the name of the desired function.

1.3.2 `from ... import`

There is another way to import functions. For example, you may wish access to all of the functions in NumPy without having to type the “`numpy.`” prefix before them. Try this:

```
from numpy import *
sqrt(2)
```

This is convenient, but it can lead to trouble when you want to use two different modules simultaneously. There is a module called `math` that also has a `sqrt` function. If you import all of the functions from `math` and `numpy`, which one gets called when you type `sqrt(2)`? (This is important when you are working with arrays of numbers.) To keep things straight, it is usually best to avoid the “`from module import *`” command. Instead, import a module and explicitly call `numpy.sqrt` or `math.sqrt` as appropriate. However, there is a middle ground. You can give a module any *nickname* you want. Try this:

```
import numpy as np
np.sqrt(2)
```


Now we can save typing and avoid confusion when functions from different modules have the same name.

There may be times when you only want a specific function, not a whole library of functions. You can ask for specific functions by name:

```
from numpy import sqrt, exp
sqrt(2)
exp(3)
```

We now have just two functions from the NumPy module, which can be accessed without the “`numpy.`” prefix. Notice the similarity with the “`from numpy import *`” command. The asterisk is a “wildcard” that tells the import command to grab everything.

One more useful variant of importing allows you to give the function you import a custom nickname:

```
from numpy.random import random as rand
rand()
```

We now have a `random` number generator with the convenient nickname `rand`.

This example also illustrates a module within a module: `numpy` contains the module `numpy.random`, which in turn contains the function `numpy.random.random`. When we typed `import numpy`, we imported many such subsidiary modules. Instead, we can import just one function by using `from` and providing a precise specification of the function we want, where to find it, and what to call it.

1.3.3 NumPy and PyPlot

The two modules we will use most often are called NumPy and PyPlot. NumPy provides the numerical tools we need to generate and analyze data, and PyPlot provides the tools we need to visualize data. PyPlot is a subset of the much larger Matplotlib library. From now on, we will assume that you have issued the following commands:

```
import numpy as np
import matplotlib.pyplot as plt
```

This can also be accomplished with the single command

```
import numpy as np, matplotlib.pyplot as plt
```

You should execute these commands at the start of every session. You should also add these lines at the beginning of any scripts that you write. You will also need to reimport both modules each time you use the `%reset` command.

Give the `%reset` command, then try importing these modules now. Explore some of the functions available from NumPy and PyPlot. You can get information about any of them by using `help()` or any of the procedures described in Section 1.2.3. You will probably find the NumPy help files considerably more informative than those for the built-in Python functions. They often include examples that you can try at the command prompt.

Now that we have these collections of tools at our disposal, let’s see what we can do with them.

1.4 PYTHON EXPRESSIONS

The Python language has a **syntax**—a set of rules for constructing expressions and statements. In this section, we will look at some simple expressions to get an idea of how to communicate with Python. The basic building blocks of expressions are literals, variable names, operators, and functions.

1.4.1 Numbers

You can enter explicit numerical values (numeric **literals**) in various ways:

- 123 and 1.23 mean what you might expect. When entering a large number, however, don't separate groups of digits by commas. (Don't type 1,000,000 if you mean a million.)
- 2.3e5 is convenient shorthand for $2.3 \cdot 10^5$.
- 2+3j represents the complex number $2 + 3\sqrt{-1}$. (Engineers may find the name j for $\sqrt{-1}$ familiar; mathematicians and physicists will have to adjust to Python's convention.)

Python stores numbers internally in several different formats. However, it will usually convert from one type to another when necessary. Beginners generally don't need to think about this. Just be aware that Python sometimes requires an integer. Even if a value has no fractional part, Python may not interpret it as an integer (for example, `a=1.0`). If you need to force a value to be an integer (for example, when indicating an entry in a list), you can use the functions `int` or `round`.

1.4.2 Arithmetic operations and predefined functions

Python includes basic arithmetic operators, for example, +, -, *, / (multiplication), / (division), and ** (exponentiation).

*Python uses two asterisks, **, to denote raising a number to a power.*

For example, `a**2` means “`a` squared.” (The notation `a^2` is used by some other math software but means something quite different to Python.)

Unlike standard mathematics notation, you must include multiplication signs. Try typing

```
(2)(3)
a = 2; a(3)
3a
3 a
```

Each command produces an error message. None, however, generates a message like, “`You forgot a '*'!`” Python used its evaluation rules, and these expressions didn't make sense. Python doesn't know what you were trying to express, so it can't tell you exactly what is wrong. *Study these error messages*; you'll probably see them again. See Appendix D for a description of these and other common errors.

Arithmetic operations have the usual precedence (ordering).

You can use parentheses to override operator precedence.

Unlike math textbooks, Python recognizes only parentheses (round brackets) for ordering operations. Square and curly brackets are reserved for other purposes. We have already seen that parentheses can

also have another meaning (enclosing the arguments of a function). Yet another meaning will appear later: specifying a tuple. Python uses context to figure out which meaning to use.

For example, if you want to use the number $\frac{1}{2\pi}$, you might type `1/2*np.pi`. (Basic Python does not know the value of π , but NumPy does.) Try it. What goes wrong, and why? You can fix the expression by inserting parentheses. Later we'll meet other kinds of operators such as comparisons and logical operations. They, too, have a precedence ordering, which you may not wish to memorize. Instead, use parentheses liberally to specify precisely what you mean.

To get used to Python arithmetic operations, figure out what famous math problem these lines solve, and check that Python got it right:

```
a, b, c = 1, -1, -2
(-b + np.sqrt(b**2 - 4*a*c))/(2*a)
```

Recall that `np.sqrt` is the name of a *function* that Python does not recognize when it launches, but that becomes available once we import the NumPy module. When Python encounters the expression in the second line, it does the following:

1. Evaluate the **argument** of the `np.sqrt` function—that is, everything inside the pair of parentheses that follows the function name—by substituting values for variables and evaluating arithmetic operations. (The argument may itself contain functions.)
2. Interrupt evaluation of the expression and execute a piece of code named `np.sqrt`, handing that code the result found in step 1.
3. Substitute the value returned by `np.sqrt` into the expression.
4. Finish evaluating the expression as usual.

How do you know what functions are available for you? See Section 1.2.3 above: Type `dir(np)` and `dir(__builtins__)` at the IPython console prompt.

A few symbols in Python and NumPy are predefined. These do not require any arguments or parentheses. Try `np.pi` (the constant π), `np.e` (the base of natural logarithms e), and `1j` (the constant $\sqrt{-1}$). NumPy also provides the standard trig functions, but be alert when using them:

The trig functions `np.sin`, `np.cos`, and `np.tan` all treat their arguments as angles expressed in radians.

1.4.3 Good practice: Variable names

Note that Python offers you no protection against accidentally changing the value of a symbol: If you say `np.pi=22/7`, then until you change it or reset Python, `np.pi` will have that value. It is even possible to create a variable whose name supplants a built-in function, for example, `round=3`.¹⁶ This illustrates another good reason for using the “`import numpy as np`” command instead of the “`from numpy import *`” command: You are quite unlikely to use the “`np.`” prefix and name your own variables `np.pi` or `np.e`. Those variables retain their standard values no matter how you define pi and e.

When your code gets long, you may inadvertently reuse variable names. If you assign a variable with a generic name like `x` in the beginning, you may later choose the same name for some completely different

¹⁶ This can be undone by deleting your version of `round`: Type `del(round)`. Python will revert to its built-in definition.

purpose. Later still, you will want the original `x`, having forgotten about the new one. Python will have overwritten the value you wanted, and puzzling behavior will ensue. You have a **name collision**.

It's good practice to use longer, more meaningful names for variables. They take longer to type, but they help avoid name collisions and make your code easier to read. Perhaps the first variable you were planning to call `x` could instead be called `index`, because it indexes a list. Perhaps the second variable you were planning to call `x` could logically be called `total`. Later, when you ask for `index`, there will be no problem.

Keep in mind, however, that "meaningful" in this context implies "meaningful to a human reader." Python itself pays no attention to the meaning of your variable names; for example, naming a variable `filename` will not tell Python how to use that variable.

Variable names are case sensitive, and most predefined names are lowercase. Thus, you can avoid some name collisions by including capital letters in variable or function names you define.

Blank spaces and periods are not allowed in variable names. Some coders use capitalization in the middle of variable names ("camel case") to denote word boundaries—for example, `whichItem`. Others use the underscore ("snake case") instead, as in `which_item`. Variable names may also contain digits (`myCount2`), but they must start with a letter.¹⁷

Some variable names are forbidden. Python won't let you name variables `if`, `for`, `lambda`, or a handful of other **reserved words**. You can find them with a web search for `python reserved words`.

1.4.4 More about functions

You may be accustomed to thinking of a function, for example, square root, as a machine that takes one number as input (its argument) and returns another number (its result) as output. Some Python functions do have this character, but Python has a much broader notion of function. Here are some illustrations. (Some involve functions that we have not seen yet.)

- A function may take a single argument, multiple arguments separated by commas, or none at all.
- A function may allow a *variable* number of arguments, and behave differently depending on how many you supply. For example, we will see functions that allow you to specify options by using *keyword arguments*. Each function's help text will describe the allowed ways of using it.
- A function may also *return* more than one value. The number of values returned can even vary depending on the arguments you supply. You can capture the returned values by using a special kind of assignment statement.¹⁸
- A function may change your computer's state in ways other than by returning a result. For example, `plt.savefig` saves a plot to a file on your computer's hard drive. Other possible side effects include writing text into the IPython console: `print('hello')`.

If you use a function name without parentheses, you are referring to the function instead of evaluating it. In mathematics, f is a function; $f(2)$ is the value of the function when its argument is 2. Type `np.sqrt` with no parentheses at the IPython command line to see how Python handles function names.

When evaluating a function, always include parentheses—even if there are no arguments.

¹⁷ Strictly speaking, a name may also begin with the underscore character, but normally such names are reserved for Python's internal use.

¹⁸ Section 6.1.4 (page 79) discusses the values returned by functions in more detail. ¹² More precisely, a Python function always returns a single object. However, that object may be a tuple that contains several items.

If a function accepts two or more arguments, how does it know which is which? In mathematical notation, the *order* of arguments conveys this information. For example, if we define $f(x, y) = x e^{-y}$, then later $f(2, 6)$ means $2 \cdot e^{-6}$: The first given value (2) gets substituted for the first named variable in the definition (x), and so on. This **positional argument** scheme is also the standard one used by Python. But when a function accepts many arguments, relying on order can be annoying and prone to error. For this reason, Python has an alternative approach called **keyword arguments**. For example,

```
f(y=6, x=2)
```

instructs Python to execute a function named `f`, initializing a variable named `y` with the value 6 and another named `x` with the value 2. You need not adhere to any particular order in giving keyword arguments. (However, keyword arguments must follow all positional arguments and you must use their correct names, which you can learn from the function's documentation.) Many functions will let you omit specifying values for some or all of their keyword arguments; if you omit them, the function supplies default values. Keyword arguments will be discussed further in Section 6.1.3.

You now know enough Python to do simple calculations. Try the examples from this chapter and play around on your own. In the next chapter, we will explore how to write simple programs in Python.

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CHAPTER 2

Organizing Data

Heisenbug: *A computer bug that disappears or alters its characteristics when an attempt is made to study it.*

— Jargon File 4.4.7

Much of the power of computation comes from the ability to do repetitive tasks extremely rapidly. You need to understand how to formulate instructions for this sort of task, so that your electronic assistant can do all the steps without your supervision. An important part of the process is organizing the data. This chapter describes several Python *data structures* that are useful in scientific computing.

2.1 OBJECTS AND THEIR METHODS

In Python, everything is an **object**. An object is a combination of data and functions. Even simple things like integers are objects in Python. (Type `dir(1)` to see all the data and functions associated with what you might have thought was “just a number.”) When processing an assignment statement, Python attaches, or **binds**, a name (a symbol like `x` or `filename`) to an object.¹ Later, you can refer to the object by this name, or you can reassign the name to a different object. Names used in this way are also called **variables**.

Let’s look at a few examples to see how this works.

When you type `i=5280`, Python begins by evaluating the right-hand side of the assignment statement. The only thing it finds is a numeric **literal**, that is, an expression that needs no evaluation. (Thus, `1+1` is an expression, but it is not a literal. In contrast, `5280` is a literal because its value is `5280`.) Continuing with the assignment, Python creates an object of type **int** to store the number `5280`. (The value of an **int** object is restricted to be an **integer**.) To complete the assignment, Python binds the name `i` to the new object. If no variable with the name `i` exists, then Python will create one. If `i` already exists, Python reassigns it to the new integer object.

When you type `f=2.5`, or `f=2.5e30`, Python creates a different type of object, called a **float**, which is short for **floating-point number**. As in scientific notation, the decimal point can “float” to give a constant number of significant figures, regardless of how large or small the number is.

Similarly, typing `s='Hello'` creates a **str** object, whose value is the **string literal** `'Hello'`, then binds the name `s` to that object. (Section 2.3 will discuss strings.) It does not matter if `s` was previously bound to a **float** or some other object. Types belong to objects. A variable can be assigned to any object.

Writing `L=[1, 2, 3]` creates a **list** object whose value is a sequence of three **int** objects. (Section 2.2

will discuss lists.)

¹  Appendix F describes in greater detail how Python handles assignment statements.

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There is more to an object than just its value. In general, objects consist of both **data** (which are often numbers) and **methods** (a particular kind of functions). The value of an object is part of its data. A method of an object is a specialized function that can act on that object's data. A method may also accept additional arguments. To access a method, you have to provide the name of the object, followed by a period, the name of the method, and a pair of parentheses. (If there additional arguments, they go inside the parentheses.) Try the following with the `float` object `f`, the `str` object `s`, and the `list` object `L` introduced above:

1. `f.is_integer()`: Every `float` object has a method called `is_integer` that determines whether its value is equivalent to an integer. The method returns `True` if the fractional part of the value is zero. Because the value of `f` is 2.5, the method returns the value `False`. The function `f.is_integer` does not require any additional arguments, but we must nevertheless place an empty pair of parentheses after its name.
2. `s.swapcase()`: Every `str` object has a method called `swapcase` that returns a new string whose value is the original string with the case of every letter reversed (upper↔lower).
3. `L.reverse()`: Every `list` object has a method called `reverse` that does not return a value but does modify the value of `L`. The output of `print(L)` shows that the order of the list has been reversed.
4. `L.pop(0)`: Every list also has a method called `pop` that will return the item at a specified location and remove it from the list. Thus, this method returns a value *and* modifies the object's data. When called with no argument, `L.pop()` will remove the last item in the list.

Every object possesses its own data and methods. Even literals have standard methods appropriate to their type: Try `'Hello'.swapcase()`, or `(5.0).is_integer()`. You can see all the methods associated with an object by using the `dir` function.

A method can use its parent object's data (examples 1–4 above). It can modify that data (examples 3, 4) or not (examples 1, 2). It can return a value (examples 1, 2, 4) or not (example 3). It can accept additional arguments in parentheses (example 4) or not (examples 1–3). Each method's documentation describes its behavior and usage.

There are some objects in Python whose values are fixed once the object is created. They are called **immutable objects**.

The methods of an immutable object do not change its value.

String and numeric objects are immutable, which is why examples 1 and 2 above do not change their values.² Lists are **mutable**: Their methods can modify their data. Examples 3 and 4 illustrated this. Shortly, we'll introduce NumPy arrays, which are also mutable.

Try creating all the example objects mentioned in this section and see how they appear in Spyder's Variable Explorer pane. Experiment with some of their methods.

Many objects have data fields (sometimes called "attributes" or "properties") that describe properties of the object other than its value. These are stored as part of the object and require no computation. Python just looks up the requested information and returns it. The syntax for accessing such properties is `object.property`. For example, try `q=1+3j` followed by `q.imag`. You should see the `imaginary` part of the complex number $1 + 3\sqrt{-1}$. No parentheses are needed because a data field is not a function. The

expression starts with the name of the broader thing (here `q`), followed by a period and the name of a specific thing (here `imag`).

² The method `swapcase` returns a modified copy of the original string; it does not affect the original string.

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You can create your own objects, and give them whatever data and methods you like, but that goes beyond the scope of this chapter. To learn more, see Chapter 10 or search the web for “python classes.”

2.2 LISTS, TUPLES, AND ARRAYS

Much of the power of computer programming comes when we handle numbers in batches. A batch of numbers could represent a single mathematical object such as a force vector, or it could be a set of points at which you wish to evaluate a function. To process a batch of numbers, we first need to collect them into a single data structure. The most convenient Python data structure for our purposes is the NumPy array, described below. Lists and tuples are also useful.

2.2.1 Creating a list or tuple

Python comes with a built-in `list` object type. Any set of objects enclosed by square brackets and separated by commas creates a list.³

```
L = [1, 'a', max, 3 + 4j, 'Hello, world!']
```

A list can contain anything; in the example just given, `max` is the name of a function.

A **tuple** is similar to a list, but immutable. To create a `tuple` object, type `t=(2,3,4)`. Note the use of round parentheses, `()`, in contrast to the square brackets, `[]`, of a list. Python knows you want a tuple because it scans the text between the parentheses and finds a comma. Without any commas, parentheses are interpreted as specifying the order of operations.⁴ Like a list, a tuple consists of an ordered sequence of objects. Unlike a list, however, a tuple is immutable. Its elements cannot be reassigned, nor can their order be modified. We will use tuples to specify the shape of an array in Section 2.2.2 and later sections. In addition, a function may return a tuple containing several objects.

2.2.2 NumPy arrays

A list is a sequence (ordered collection) of objects, endowed with methods that can perform certain operations on its contents like searching and counting. You can do numerical calculations with lists; however, a `list` object is generally *not* the most convenient data type for such work. What we usually want is a *numerical array*—a grid of numbers, all of the same type. The NumPy module provides a class of array objects ideally suited to our needs. Knowing that all of the elements in an array are numbers of the same type allows NumPy to do efficient calculations on an entire array. Because Python lists can contain any mix of object types, processing their elements is far less efficient.

*A Python list is **not** the same as a NumPy array.*

We will almost always use NumPy arrays.

You can create a one-dimensional array by using the command

```
a = np.zeros(4)
```

³ A list can contain just one object, but note that `2.71`, a `float` object, is not the same thing as `[2.71]`, a list containing one `float` object. A list can even contain *no* objects: `L=[]`.

⁴ If you want a tuple with just one element, write `(0,)`.

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The function `np.zeros` requires one argument that describes the shape of the array; in this case, the integer `4` specifies an array with four elements. It sets all the entries to zero.

In mathematics, we often need arrays of numbers with two or more dimensions. Try

```
a = np.zeros( (3, 5) )
```

and see what you get. Here `np.zeros` again accepts a single argument. In this case, it is the tuple `(3, 5)`. The function creates an array with 15 entries, all equal to zero, arranged in three rows and five columns. The name `a` now refers to a `ndarray` object (an “**n**-dimensional NumPy `array`”) that conforms to the mathematical notion of a 3×5 matrix.

Your
Turn
2A

Create `a` by using the preceding code. Then, find it in the Variable Explorer. Double-click its value to look inside the array. Repeat with the function `np.ones` in place of `np.zeros` and see what you get. Finally, try `np.eye(3)`.

A *row vector* is a special case of a two-dimensional array with just one row: `np.zeros((1, 5))`. Similarly, a *column vector* is a two-dimensional array with just one column: `np.zeros((3, 1))`. Python will give you exactly what you ask for, so be aware of what you are requesting:

If you want a column vector of N zeros, use `np.zeros((N, 1))`, not `np.zeros(N)`. If you want a row vector of N zeros, use `np.zeros((1, N))`.

The functions `np.ones` and `np.random.random` can also create two-dimensional arrays, using the same syntax.

Python can report how big an array is. After setting up `a`, use the command `np.size(a)` and see what you get. Also try `np.shape(a)`. This information is also displayed in the Variable Explorer.

A NumPy array is an object with its own data. For example, it can report its own size and shape (`a.shape` and `a.size`). A NumPy array also has methods for acting on its data. Try `a.sum()`, `a.mean()`, and `a.std()`. (Of course, these methods are more useful when the array contains something other than zeros!)

Compare the output of `np.zeros(3)`, `np.zeros((1, 3))`, and `np.zeros((3, 1))`. Note, in particular, the shape of each. NumPy regards the first as neither a row nor a column vector. It is a one-dimensional array—an array with only one index. In some cases, NumPy will treat all three types of arrays the same. However, in certain matrix and vector operations, the shape of the array is important. Be certain you have arrays of the shape you need when performing these operations. You can reshape a one-dimensional array into a column vector or row vector if necessary. (See Section 2.2.9.)

Python can also handle arrays with more than two dimensions. Try

```
A = np.zeros( (2, 3, 4) )
B = np.ones( (2, 3, 4, 3) )
```

and inspect the resulting arrays.

Don't confuse the concept of "three-dimensional array" with that of "three-dimensional vector." The variable `A` defined earlier is a three-dimensional array; it could represent grid points filling a volume of space, each holding a single quantity (a *scalar*). In contrast, a three-dimensional vector is just a set of

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three numbers (the components of the vector); it could be represented by an array like `np.ones(3)` or `np.ones((3,1))`.⁵

2.2.3 Filling an array with values

Perhaps you'd like to set up an array with more interesting values. The command

```
a = [2.71, 3.14, 3000]
```

creates a Python list—not a NumPy array—with the specified values. It contains the values we want, but we cannot perform the operations we would like with the list. To create a NumPy array from these values, we can call the function `np.array` on the list:

```
a = np.array([2.71, 3.14, 3000])
```

Making each element of the list its own list—that is, enclosing each entry in square brackets—returns a column vector, a two-dimensional array with three rows each consisting of a single column:

```
a = np.array([[2.71], [3.14], [3000]])
```

Your
Turn
2B

Try

```
a = np.array([[2, 3, 5], [7, 11, 13]])
```

and explain the result.

Usually you don't want to specify each entry in an array explicitly. For example, we frequently want to create a NumPy array of evenly spaced values over some range. NumPy provides two useful functions to do this: `np.arange` and `np.linspace`.

The function `np.arange(M,N)` creates a one-dimensional array with M as its first entry, M+1 as the second, and so on, stopping just *before* it reaches a value that equals or exceeds N.

Example: Try the following commands. Inspect and describe the resulting arrays.

```
a = np.arange(1, 10)
b = np.arange(5)
c = np.arange(2.1, 5.4, 0.1)
```

The last command creates a series, but each entry is greater than its predecessor by 0.1, not by the default increment of 1.

The `arange` syntax is `np.arange(start, end, increment)`. The increment and starting value are optional. The default start value is 0; the default increment is 1. The end value is not included in the array.

Type `help(np.arange)` at the command prompt to learn more.

⁵ A three-dimensional grid of three-dimensional vectors could be represented by a *four*-dimensional NumPy array!

The function `np.linspace(A, B, N)` does a similar job. It creates a one-dimensional array with exactly N evenly spaced entries. The first entry equals A . Unlike the `np.arange` function, however, the last entry equals B exactly. You don't get to specify the spacing; Python determines what is needed to cover the range from A to B with N equally spaced points.

*The `linspace` syntax is `np.linspace(start, end, num_points)`.
The end value is included in the array.*

Your Turn 2C

a. Try

```
a = np.arange(0, 10, 2)
b = np.linspace(0, 10, 6)
```

and explain the results. See if you can use `np.arange` and `np.linspace` to create identical arrays. [Hint: What happens when you add the increment to the end value in `np.arange`?]

b. Now try

```
a = np.arange(0, 10, 1.5)
b = np.linspace(0, 10, 7)
```

Describe how `a` and `b` differ. Explain why, and decide which form you should use if you want to evaluate a function over the range 0 to 10.

When creating a series to compute the values of a function over a range, `np.linspace` is the appropriate function. It allows you to specify the start and end of the range (and the number of points in the series). However, when the exact spacing between points is important, use `np.arange` instead.

If you want to control both the end point and the spacing, the following construction will do the job:

```
x_min = 0
x_max = 10
dx = 0.1
x_array = np.arange(x_min, x_max + dx, dx)
```

Python also has a built-in function called `range`, but it does not create a numerical array of values. Instead, it creates an object that returns a sequence of values, one at a time. This makes `range` useful for loops, as described in Section 3.1, but it should not be used in place of an array.

2.2.4 Concatenation of arrays

NumPy offers two useful methods for building up an array from smaller ones. Each takes a single argument, which is a list (or a tuple) whose entries are the arrays to be combined:

- **`np.hstack` (horizontal stack):** The resulting array has the same number of rows as the original arrays. The arrays to be stacked must have the same number of rows.
- **`np.vstack` (vertical stack):** The resulting array has the same number of columns as the original arrays. The arrays to be stacked must have the same number of columns.

Try these examples, and describe the results:

```
a = np.zeros( (2, 3) )
b = np.ones( (2, 3) )
h = np.hstack( [a, b] )
v = np.vstack( [a, b] )
```

Look at the contents of `h` and `v`, and compare their shapes with those of `a` and `b`.

2.2.5 Accessing array elements

Once you have created an array, list, or tuple, you can access each of its entries individually. Try the following code at the command prompt:

```
A = np.array( [2, 4, 5] )
A[0]
A[1] = 100
print(A)
```

Array indices (offsets) are enclosed in square brackets, not parentheses.

Using round (ordinary) parentheses to request array elements is a common error when learning Python.

The third line above changes just one entry in the array, but it may not be the entry you expect. In Python, the indices of lists, tuples, arrays, and strings all start with 0. Thus, `A[1]` is the *second* element of `A`. This is not the convention that math texts use for vectors and matrices.⁶ (The same indexing scheme can access individual characters in strings.)

Indices in Python start at 0. If A is a one-dimensional array with N elements, the first element is A[0] and the Nth element is A[N-1]. Asking for element A[N] will result in an error.

A math text would refer to `A[N-1]` as A_N . The notation A_{-1} does not appear in most math texts, but Python interprets a negative index as an offset from the *end* of a list or array. More precisely, `A[-1]` will access the last entry of an array, and `A[-n]` refers to the *n*th element from the end of an array.

To understand how to access elements in a two-dimensional array, try the following:

```
A = np.array( [ [2, 3, 5], [7, 11, 13] ] )
A[0]
A[0][1]
A[1][2] = 999
```

In the second line above, `A[0]` returns an array whose elements are `[2, 3, 5]` (the first row). In the third line, `A[0][1]` then asks for the *second* item in *that* array, which is the number 3. NumPy arrays also understand an abbreviated indexing scheme:

```
A = np.array( [ [2, 3, 5], [7, 11, 13] ] )
A[0, 1]
A[1, 2] = 999
```

⁶  The convention in Python is inherited from another programming language called C. In that language, the name of an array is linked to the location in memory where its first value is stored. An index is interpreted as an offset. Thus, to get the *first* entry in the array, we need an offset of *zero*: `A[0]`.

Both `A[i][k]` and `A[i,k]` return the entry at the intersection of row `i+1` and column `k+1`. A mathematics text would call this entry $A_{i+1,k+1}$. Again, Python's behavior is easier to understand if you think in terms of offsets: `A[i,k]` is the entry `i` steps down and `k` steps to the right from the upper-left corner of the array.

2.2.6 Arrays and assignments

An assignment statement for a variable, such as `f=2.5`, looks similar to an assignment statement for an element of an array, such as `A[1]=2.5`. However, these statements are handled differently in Python, and it is important to understand the distinction.

When we access and assign elements of an array, we are using methods of an `ndarray` object to view and modify its data.⁷ In contrast, when we assign a value to a variable, as in `f=2.5`, Python binds the variable name to a new object. There is little difference in behavior until we have two names bound to the same object.

Suppose that we create two variables that point to the same `float` object and two variables that point to the same `ndarray` object:

```
f = 2.5
g = f
A = np.zeros(3)
B = A
```

Inspect all four variables, `A`, `B`, `f`, and `g`, before and after these commands:

```
g = 3.5
A[0] = 1
B[1] = 3
```

The first statement has no effect on `f`, even though the value of `g` is changed. However, `A` and `B` remain equal to each other. The reason is that `A` and `B` are bound to the same `ndarray` object, and each used a method of that object to modify its data.

If multiple variables are bound to the same array, they can all use its methods to modify its data.

This includes assignment of individual elements.

Lists behave in a similar way. (Tuples are immutable objects that do not allow reassignment of their elements, so `A[0]=1` results in an error if `A` is a tuple.)

There is another important difference between variable assignment and array assignment. If you type `A=1`, Python does not raise an error, even if `A` was previously undefined. However, if you try `A[0]=1` before defining `A`, Python returns an error message. The reason is that `A[0]=1` is attempting to use a method of an object named `A` to modify its data. Since no such object exists, Python cannot proceed. If you need to set the entries of an array one by one, you should first create an array of the appropriate size, for example, `A=np.zeros(100)`.

⁷ The expression `A[1]` is shorthand for a method called `A.__getitem__(1)`, and `A[1]=2.5` calls the method `A.__setitem__(1,2.5)`.

2.2.7 Slicing

Often you will wish to extract more than one element from an array. For example, you may wish to examine the first ten elements of an array or the 17th column of a matrix. These operations are possible using a technique called **slicing**.

The syntax for slicing is `a[start:end:stride]`.

In Python, a colon indicates slicing when it is part of an index. Thus, `a[start:end:stride]` returns a new array whose entries are

```
a[start], a[start + stride], a[start + 2*stride], ..., a[start + M*stride]
```

where M is the largest integer for which $\text{start} + M*\text{stride} < \text{end}$. The stride comes last. If it is omitted, the default is 1. If `start` or `end` are omitted, their defaults are the first and last indices, respectively. If the stride is omitted, the second colon may also be omitted.

If you want to see the first ten entries in `a`, you could use the expression `a[0:10:1]`, or more concisely `a[:10]`. The same syntax can be used to slice along each dimension of a multidimensional array. For example, to get a slice of the third column of a matrix, you could use `A[start:end:stride,2]`.

Suppose that you have been given experimental data in an array `A` with two columns. For your analysis, you need to split the two columns of data into separate arrays. If you don't know the number of rows in `A`, you could start by finding it, and then assign the slices to separate variables:

```
N = np.size(A, 0)
x = A[0:N:1, 0]
y = A[0:N:1, 1]
```

A colon all by itself represents every allowed value of an index (that is, `start=0, end=-1, stride=1`). We can use this shortcut to specify our slices more concisely:

```
x = A[:, 0]
y = A[:, 1]
```

Your
Turn
2D

Try the following commands to familiarize yourself with slicing:

```
a = np.arange(20)
a[:]
a[::]
a[5:15]
a[5:15:3]
a[5::]
a[:5:]
a[::5]
```

Explain the output you get from each line.

Can you construct a slicing operation that returns only the odd entries of `a`?

Negative index values can also be used in slicing. This can be especially useful if you only want to see the last few entries in an array whose size is unknown. For example, `a[-10:]` will return the last ten

elements stored in `a` (if `a` has at least ten elements). Similarly, `a[:-10]` is a slice that stops ten elements before the end of the array—everything except the last ten elements stored in `a`.

An array slice can also appear in an assignment statement:

```
A = np.zeros(10)
A[0:3] = np.ones(3)
```

This code replaces a block of values in `A`, leaving the rest unchanged. (See Section 2.2.6.)

2.2.8 Flattening an array

An array may have any number of dimensions. In some cases, you may wish to repackage all of its values as a one-dimensional array. The function `np.ravel` does exactly this. It can be accessed as a NumPy function or an array method. NumPy arrays can also *flatten* themselves. Try the following commands, and inspect the results:

```
a = np.array( [ [1, 2], [2, 1] ] )
b = np.ravel(a)
c = a.ravel()
d = a.flatten()
```

Notice, in particular, that neither `ravel` nor `flatten` changes `a`. Each returns a one-dimensional array that contains the same elements as `a`. There is a significant difference between the methods, however. The `flatten` method returns a new, independent array; `ravel` returns an `ndarray` object with access to the *same* data as `a`, but with a different shape. To see this, compare the effects of the assignments `d[1]=11` and `b[2]=22` on all four arrays.

2.2.9 Reshaping an array

Flattening an array is just one way to change its shape. An array can be recast into any shape consistent with its number of elements by using `np.reshape` or an array's own `reshape` method. Try the following, and inspect each array:

```
a = np.arange(12)
b = np.reshape(a, (3, 4) )
c = b.reshape( (2, 6) )
d = c.reshape(2, 3, 2)
```

The `reshape` method takes a tuple of numbers, like `(3, 4)` or `(2, 3, 2)`, as its argument. (When you call the `reshape` method of an array, you do not need to enclose the new shape in parentheses.) As long as the product of these numbers is equal to the number of elements in the original array, the command will return a new array. If one of the entries in the requested shape tuple is `-1`, then `np.reshape` substitutes whatever integer will make the total number of entries match that of the input array.

Like `ravel`, these `reshape` methods return an `ndarray` object with access to the same data as `a`, but with a different shape. Modifying the elements of any of the arrays in this example will affect all four of them.⁸

⁸ `T2` Array slicing, reshaping, and NumPy's `ravel` method each return a `view` of the original array. That is, the resulting object has access to the same data as the original array. Any changes to a view change the original array—and all other views of that array. See Appendix F.

Reshaping provides a convenient way to transform any array into a row vector or column vector:

```
z = np.arange(10)
z_row = z.reshape(1, -1)
z_col = z.reshape(-1, 1)
```

(Other methods are described in Section 2.2.3.) You are allowed to pass `-1` as one (and only one) dimension when calling the `reshape` function. NumPy interprets this as a request to reshape the array, using whatever value is required for this dimension. If you supply `1` as the other dimension, NumPy will place all of the elements of the array in a single row or column.

2.2.10 Lists and arrays as indices

Python has an even more flexible method for slicing NumPy arrays: You can use a list or array where an index is expected. Using a list of integers will return an array that contains just the elements of the original array at the offsets specified by the list. Try

```
a = np.arange(10, 21)
b = [2, 4, 5]
a[b]
```

This method can be extremely useful if the list `b` was itself generated automatically. For example, `b` could be a list of time points at which we wish to sample a signal contained in `a`.

You can also use a **Boolean** array (an array whose entries are either `True` or `False`) to select entries from another array of the same shape. This technique is called **logical indexing**. Try the following:

```
a = np.arange(-10, 11)
less_than_five = (abs(a) < 5)
b = a[less_than_five]
```

The comparison in the second line returns an array with the same shape as `a`, whose entries are `True` or `False`, depending on whether the particular element in `a` satisfies the comparison.⁹ When `less_than_five` is used as an index to `a`, Python returns an array containing only those elements of `a` for which the corresponding element in `less_than_five` is `True`.

It is not necessary to create a named array to use as an index. The following line is equivalent to the last two lines of the example above:

```
b = a[abs(a) < 5]
```

2.3 STRINGS

Python can manipulate text as well as numbers. Next to an array, the second most important data structure for our purposes is the **string**. A string may contain any number of characters. You can create one with

```
s = 'Hello, world!'
print(s)
type(s)
```

⁹ This is an example of *vectorized* computation, which will be discussed in Section 3.2.1.

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Notice the output of `type(s)`.

The expression on the right side of the equals sign above is a **string literal**. The equals sign assigns this short sentence as the value of `s`. Python allows you to use either a single quote (') or a double quote ("") to define a string; however, you must begin and end the string with the same character.

A string literal starts and ends with a single quote or a double quote.

A single quote is the same as the apostrophe on your keyboard. Elsewhere on your keyboard there's a different key, the grave accent. You may be tempted to use it as a left quote, as in `string'. Python won't understand that.

If you need an apostrophe inside a string, you can enclose the whole string in double quotes: `"Let's go!"` If you need both an apostrophe *and* double quotes, you will have to use a backslash (\) before the symbol that encloses the string. Write either `"I said, \"Let's go!\""` or `'I said, "Let\'s go!"'` More generally, inside a string, the backslash is an **escape character** that means, "Interpret the next character literally." In these examples, it instructs Python that the double quotes and the apostrophe, respectively, are part of a string, not the beginning or end of a string.¹⁰

A string may contain a collection of digits that looks like a number, for example, `s='123'`. However, Python *still considers such a value to be a string*, not the number 123. Try typing

```
a = '123'  
b = a + 1
```

Python issues a **TypeError** when you try to add a number to a string. However, you can *convert* a string to a number. (This may be necessary if you are getting input from a keyboard.) Python can do sensible conversions like creating an integer from `"123"` or a **float** from `"3.14159"`, but it does not know how to directly create an integer from a string that contains a decimal point. Try the following:

```
s = '123'  
pie = '3.142'  
x = int(s) + 1  
y = float(pie) + 1  
z_bad = int(s) + int(pie)  
z_good = int(s) + int(float(pie))
```

Python does know how to "add" two strings together. Try this:

```
"2" + "2"
```

The result is not what you learned in kindergarten. Python uses addition (+) to *concatenate* (join) strings:

```
s = 'Hello, world!'  
t = 'I am Python.'  
s + t
```

Your
Turn
2E

The result of the last evaluation doesn't look quite right. Replace the last line with `s+ ' '+t` and explain the operations that lead to the output.

¹⁰ Another use of backslash is to code special characters, such as newline (\n) and tab (\t). See also page 59. You can even code

..... backslash itself via \\.

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Some Python functions require arguments that are strings. For example, graph titles and legends must be specified as strings. (See Section 4.3.) You may wish to include the value of a variable in such a string. If the value is not already a string, one option is to convert it. Just as Python can convert strings to integers and floating point numbers, it can convert numbers to strings. The built-in function `str` will try to create a string using whatever input you provide:

```
s = "Poisson distribution for $\mu$ = " + str(mu_val)
```

This string can now be used by PyPlot to generate a graph title. In this expression,

- The `+` sign joins two strings.
- The first string is a literal. It contains some special characters that PyPlot can use to produce the Greek letter μ .¹¹
- The second string is obtained from the current value of a variable named `mu_val`.

Python will display up to 16 digits when printing floating point numbers, and this is what the `str` command will generate in the example above if μ has that many digits. This may be satisfactory, but Python offers much more control over the appearance of strings. You can control how numbers appear in strings by using special commands involving the percent sign (%) or the `format` method that every string possesses. The percent sign is widely used, so we will briefly describe it below. However, the `format` method behaves more like other functions in Python, so we will discuss it first and use it throughout the remainder of the tutorial.

2.3.1 Raw strings

Strings that contain backslashes, such as \LaTeX commands or Windows filenames, are awkward to type because of all of the escape characters. Using a **raw string** can simplify things. In a raw string, there are no escape characters: A backslash is just a backslash. To create a raw string, simply precede the first quotation mark with the letter `r`:

```
# Windows paths
path1 = "C:\\Documents\\code\\data1.csv" # normal string
path2 = r"C:\\Documents\\code\\data2.csv" # raw string

# LaTeX strings
latex1 = "$\\cos \\theta = \\frac{\\sqrt{3}}{2}$" # normal string
latex2 = r"$\\cos \\theta = \\frac{\\sqrt{3}}{2}$" # raw string
```

There is no difference in the meaning of `path1` or `path2`, or between `latex1` and `latex2`, as you can verify with the `print` command. These raw strings are easier to type and less prone to errors, but this comes at the cost of flexibility. You cannot create raw strings with newlines, tabs, or other special characters unless you use one of the string formatting methods below.

2.3.2 Formatting strings with the `format()` method

Try some examples that use the `format()` method.¹²

¹¹ Python will interpret the text between dollar signs (\$...\$) as \LaTeX typesetting instructions. The first backslash escapes the second one and prevents it from being interpreted as an escape character itself.

¹² The hash symbol # in the following code snippet introduces a comment. (See Section 3.3.4.) The comment here tells you the name of the code snippet, so that you can find it in the book's online resources.


```
# string_format.py      [get code]
"The value of pi is approximately " + str(np.pi)
"The value of {} is approximately {:.5f}.".format('pi', np.pi)
s = "{1:d} plus {0:d} is {2:d}"
5 s.format(2, 4, 2 + 4)
"Every {} has its {}".format('dog', 'day', 'rose', 'thorn')
"The third element of the list is {0[2]:g}.".format(np.arange(10))
```

When evaluating a string's `format` method, Python interprets a pair of curly brackets (`{}`) as a placeholder, where a value will be inserted. The arguments of `format` are expressions whose values will be inserted at the designated points. They can be strings, numbers, lists, or more complicated expressions. These examples illustrate several properties of the `format` method:

- An empty pair of curly brackets will insert the next item in the series of expressions, as illustrated in the third line. The item can be formatted if the brackets contain a colon (`:`) followed by a formatting command. For example, “`{:.5f}`” means, “Format the current argument as a floating point number with 5 digits after the decimal point.”
- Items can also be explicitly referenced by their index in the series of expressions passed to `format`. The fourth line in the examples above defines a string with three placeholders to be filled later. It will take the second, then the first, then the third arguments of `format` and insert them at the designated locations. The syntax “`{1:d}`” means, “Insert the second argument here, and display it as a decimal (base 10) integer.” (Python can also represent integers as binary, octal, or hexadecimal by using `{:b}`, `{:o}`, and `{:x}`, respectively.)
- Line 6 shows that not all arguments need to be used.
- The last line demonstrates how you can reference an individual item of an array passed to `format`: The *replacement field* `0[2]` refers to the third element of the first argument, and the *format specifier* `:g` then instructs Python to display a number using the fewest characters possible (general format). It may choose exponential notation, and it will leave off trailing zeros in floating point numbers.

Python's `help(str)` information provides basic descriptions of the available string methods. There are also many references online if you wish to explore string processing beyond the rudimentary introduction provided here. However, it can be quicker and more fun to

Experiment in the console to discover how things work.

2.3.3 Formatting strings with %

Try the following at the command prompt to see how the `%` syntax works:

```
# string_percent.py      [get code]
"The value of pi is approximately " + str(np.pi)
"The value of %s is approximately %.5f" % ('pi', np.pi)
s = "%d plus %d is %d"
5 s % (2, 4, 2 + 4)
```

When a string literal or variable is followed by the `%` operator, Python expects that you are going to provide values to be formatted and inserted into that string. The desired insertion point(s) are indicated by additional `%` characters within the string. These can be followed by information about how each value

should be formatted, as described in Section 2.3.2. As these examples show, you can insert multiple values into a string, and each value can be the result of evaluating an expression. You must provide the values in the order they are to appear in the string.

Thus, in the third line of the example above, “%s” means, “Insert a **string** here.” Similarly, “%.5f” means, “Insert a floating point number here with 5 digits after the decimal point.” In the third line of the example, “%d” means, “Insert a **decimal** (base 10) integer here.”

In addition to formatting strings, the percent sign can also function as an arithmetic operator in Python. In an expression like $5 \% 2$, the percent sign represents the **modulo operation**. It returns the remainder upon division. (Try it.) Because of its multiple meanings, bugs that involve a percent sign can lead to puzzling behavior that is difficult to diagnose.

This chapter introduced the most common Python data structures in scientific computing. This is enough to get started with array processing and string manipulation. However, to get the most out of Python, we need to do more than enter instructions at the command line. It is time to look at the flow of operations in a chain of instructions—that is, at computer *programs*.

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CHAPTER 3

Structure and Control

When you come to a fork in the road, take it.

— Yogi Berra

The previous chapter introduced some useful structures for storing and organizing data. To utilize them effectively, we now need to automate repetitive operations on the data. This chapter describes how to group code into repeated blocks (looping) and contingent blocks (branching), and how to assemble code blocks into reusable computer programs called scripts.

3.1 LOOPS

So far, we have described Python as a glorified calculator with some string processing capabilities. However, we can continue to build on what we have learned and do increasingly complex tasks. In this section, we will explore two **control structures** that will allow us to repeat a set of operations as many times as we need: **for** loops and **while** loops.

3.1.1 for loops

Instead of solving a single quadratic equation, let's go a step further and create a *table* of solutions for various values of **a**, holding **b** and **c** fixed. To do this, we will use a **loop**. Try typing the following in the IPython console:

```
# for_loop.py      [get code]
b, c = 2, -1
for a in np.arange(-1, 2, 0.3):
    x = (-b + np.sqrt(b**2 - 4*a*c)) / (2*a)
    print("a= {:.4f}, x= {:.4f}".format(a, x))
```

(Hit <Return> on a blank line to terminate the loop and run it.)

Note the colon after the **for** statement. This is essential. It tells Python that the next lines of indented code are to be associated with the **for** loop. The colon is also used in **while** loops and **if**, **elif**, and **else** statements, which will be discussed shortly.

The keyword **for** instructs Python to perform a block of code repeatedly. It works as follows:

1. The function **np.arange(-1, 2, 0.3)** indicates a series of values with which to execute the indented block of code. Python starts with the first entry and cycles over each value in the array.
2. Python initially assigns **a** the value **-1**. Then, it evaluates a solution to the quadratic equation and prints the result. (Without the **print** statement, nothing would be displayed as the calculation proceeds.)

3. The end of the indented code block tells Python to jump back to the beginning of the loop, update the value of `a`, and execute the block *again*. Eventually, Python reaches the end of the array. Python then jumps to the next block of unindented code (in this case, nothing). We say that it “exits the loop.”

Note that `a` is an ordinary variable whose value can be accessed in calculations. It is generally not good practice to modify its value within the loop, however. When Python reaches the end of an indented block, it will continue on to the next value in the array and discard any changes you made to the value of `a`.

The example above illustrates a very important feature of Python:

*Blocks of code are defined **only** by their indentation.*

In many programming languages, special characters or commands separate blocks of code. In Java, C, or C++, the instructions in a `for` loop are enclosed in curly brackets (`{ ... }`). In other languages, the end of a block of code is designated by a keyword (for example, `end` in MATLAB). In Python, the indentation of a statement—and nothing else—determines whether it will be executed inside or outside a loop. This forces you to organize the text of your program exactly as you intend the computer to execute it. You can tell at a glance which commands are inside a loop and where a block of code ends, and your code is not cluttered with dangling curly brackets or a pyramid of `end` statements at the end of a complex loop.

How much should you indent? The only requirements in Python are the following:

- Indentation consists of blank spaces or tabs.
- Indentation must be consistent within a block. (For example, you cannot use four spaces on one line and a single tab on the next, even if the tab appears to indent the block by four spaces.)
- The indentation level must increase when starting a new block, and go back to the previous level when that block ends.

The official “Style Guide for Python Code” (PEP8) recommends using spaces instead of tabs, and using four spaces per indentation level. (See python.org/dev/peps/pep-0008.)

IPython’s command line interpreter aids you in setting up indentation. When you type the first line of a `for` loop and hit `<Return>`, the next line is automatically indented. When you write your own programs, you will have to make sure statements are indented properly. (Many semi-intelligent editing programs, including the Editor in Spyder, will also help you with indentation.)

Try this modified set of instructions and explain why the output differs from before (prior to typing the last line, hit `<Backspace/Delete>` to undo the automatic indentation):

```
b, c = 2, -1
for a in np.arange(-1, 2, 0.3):
    x = (-b + np.sqrt(b**2 - 4*a*c)) / (2*a)
    print("a= {:.4f}, x= {:.4f}".format(a, x))
```

We can tell at a glance that this version of the loop will cycle over the values of `a`, and *then* display the final values of `a` and `x` once Python exits the loop.

If the body of a loop is very brief, you can just write it on the same line as the `for` statement after the colon, and forget about indentation:

```
for i in range(1, 21): print(i, i**3)
```


3.1.2 while loops

A second example of a control structure is useful when you wish to repeat a block of code as long as some general condition holds but you do not know how many iterations will be required. A `while` loop will exit the first time its condition is `False`. For example, you could calculate solutions to a quadratic equation until the discriminant changes sign as follows:

```
# while_loop.py      [get code]
a, b, c = 2, 2, -1
while (b**2 - 4*a*c >= 0):
    x = (-b + np.sqrt(b**2 - 4*a*c)) / (2*a)
    print("a = {:.4f}, x = {:.4f}".format(a, x))
    a = a - 0.3
print("done!")
```

Note that we now need to change the value of `a` explicitly within the loop (line 6), because we are not iterating over an array. Also, be aware that a different choice of values for `a`, `b`, and `c` could result in an infinite loop. (See Section 3.1.4 below.)

As with a `for` loop, the indentation of statements tells Python which commands to execute inside a `while` loop; it also tells Python at what point to pick up again after the loop has completed. In the example above, the code prints out a short message after finishing the loop.

3.1.3 Very long loops

Some calculations take a long time to complete. While you're waiting, you may be wondering whether your code is really working. It's a good idea to make your code provide updates on its progress. If you have a loop like `for ii in range(10**6):`, then you could insert the following line:

```
if ii % 10**5 == 0: print("{:.0f} percent complete".format( 100*ii/10**6 ))
```

Here the percent sign indicates the arithmetic operation of remainder (Section 2.3.3).

3.1.4 Infinite loops

When working with loops, there is the danger of entering an **infinite loop** that will never terminate. This is usually a bug, and it is useful to know how to halt a program with an infinite loop without quitting Python itself.

The easiest way to halt a Python program is to issue a **KeyboardInterrupt** by typing `<Ctrl-C>`. This will work on most programs and commands. Try it now. At the IPython command prompt, type

```
while True: print("Here we go again ...")
```

When you hit `<Return>`, Python will enter an infinite loop. Halt this loop by typing `<Ctrl-C>`.¹

`<Ctrl-C>` can also be used to halt commands or scripts that are taking too long, or lengthy calculations that you realize are using the wrong input. Just click in the IPython console pane, and type `<Ctrl-C>`. There is also a red `STOP` button in the upper-right corner of the IPython console that

¹ If you are using a Jupyter notebook, `<Ctrl-C>` will not work. You must use the “Interrupt Kernel” button (stop button) or the three-key sequence, `<Esc>, <I>, <I>`.

will issue a **KeyboardInterrupt** from within Spyder (see Figure 1.1, page 7). Run the infinite loop command again, and halt it with the **STOP** button.

It may take a while for Python to respond, depending on what it is doing, but <Ctrl-C> or the **STOP** button will usually halt a program or command. However, if Python is completely unresponsive, you may have to exit Spyder or force it to quit with the **Restart kernel** option in the **OPTIONS** menu to the right of the **STOP** button on the IPython console pane (again see Figure 1.1). If this happens, you could lose everything you have not saved, so be sure to save your work frequently.

IPython keeps a record of the commands you entered. If you need to recover your work, or if you want to copy a useful series of commands into a separate file for later use, you can use the IPython magic command **%history**. By default, this will display all of the commands you have typed in the current session. This is not very helpful if you are attempting to recreate the state of affairs before you had to restart the kernel or restart Spyder. However, you can provide an optional argument to specify how far you would like to look back: **%history -1 100** will display the last 100 commands, spanning multiple IPython sessions if necessary. You can select and copy portions of this history to the clipboard in the usual fashion: highlight with the mouse, and then type <Cmd-C> or right-click with the mouse to copy. You can paste directly from the clipboard and run the commands by using the IPython magic command **%paste**. Your session history can be useful in recovering your work, and it can provide a starting point for Python scripts, which we discuss in the next chapter.

3.2 ARRAY OPERATIONS

One reason to use arrays is that NumPy has a very concise syntax for handling repetitive operations on arrays. NumPy can directly calculate the square root of every element in an array or the sum of each column in a matrix much more rapidly than a **for** loop designed to do the same thing. Applying an operation to an entire array instead a single number (scalar) is called **vectorization**. An operation that combines elements of an array to create a smaller array—like summing the columns of a matrix—is called **array reduction**. Let's look at examples of both of these operations.

3.2.1 Vectorizing math

We can calculate the solutions to the quadratic equation in Section 3.1.1 by using the following code:

```
# vectorize.py      [get code]
b, c = 2, -1
a = np.arange(-1, 2, 0.3)
(-b + np.sqrt(b**2 - 4*a*c)) / (2*a)
```

We type the last command exactly as we would for a single value of **a**, but Python evaluates the operation for every element of **a** and returns the results in an array.

To evaluate the expression on the fourth line, Python essentially carries out the following steps:²

1. Python starts with the innermost subexpression, first calculating **b**2** and saving the result. In evaluating **4*a*c**, it notices that the array **a** is to be multiplied by 4. Python interprets this as a request to multiply *each element* by 4. This is the standard interpretation in math: When you multiply a vector

² Python may actually use a more efficient procedure.

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3.2 Array operations 39

by a scalar, each component is multiplied by that quantity. Similarly, Python then multiplies the resulting array by `c`.

2. After evaluating `4*a*c`, Python sees that you've asked it to combine this result with the single quantity `b**2`. *Unlike* standard math usage (where it does not make sense to add a vector and a scalar together), Python interprets this as a request to create a new array, in which the entry indexed by k is equal to `b**2 - 4*a[k]*c`. That is, addition and subtraction of an array and a scalar are also evaluated item by item.
3. Python then passes the array generated in step 2 to the `np.sqrt` function as an argument. NumPy's square root function will accept a single number, but like many functions in the NumPy module (including `sin`, `cos`, and `exp`), it can also act on an array. The function acts item by item on each entry and returns a new array containing the results.³
4. Python adds `-b` to the array from 3 using the same rule as in 2.
5. Python divides the array from 4 by `2*a`. This operation involves two arrays and will be discussed in a moment. It is also evaluated item by item.

The code in `vectorize.py` is a *vectorized* form of the earlier `for` loop. Vectorized code often runs much faster than equivalent code written with explicit `for` loops, because clever programmers have optimized NumPy functions to run "behind the scenes" with compiled programs written in C and FORTRAN, avoiding the usual overhead of Python's interpreter.

Vectorization can also help you write clearer code. Long, rambling code can be hard to read, making it difficult to spot bugs. Of course, extremely dense code is also hard to read. Your coding style will evolve as you get more experienced. Certainly if a code runs fast enough with `for` loops, then there's no need to go back and vectorize it.

We can now finish explaining the example at the start of this section. The variable `a` is an array; therefore, `-b + np.sqrt(b**2-4*a*c)` and `2*a` are also arrays. When two NumPy arrays of the same shape are joined by `+`, `-`, `*`, `/`, or `**`, Python performs the operation on each pair of corresponding elements, and returns the results in a new array of the same shape.

Not every mathematical operation on an array will act item by item, but most common operations do. Suppose that you wanted to graph the function $y = x^2$. You could set up an array of x values by `x=np.arange(21)`. Then, `y=x**2` or `y=x*x` gives you what you want.

Your
Turn
3A

- a. You may wish to evaluate the function $y = e^{-(x^2)}$ over a range of x values, for example, to graph a normal probability distribution. Figure out how to code this in vectorized notation.
- b. We often wish to evaluate the function $e^{-\mu} \mu^n / (n!)$ over the integer values $n = 0, 1, \dots, N$. Here, the exclamation point denotes the factorial function. Figure out how to code this in vectorized notation for $N = 10$ and $\mu = 2$. (You may want to import the `factorial` function from SciPy's collection of special functions, `scipy.special`.)

The item-by-item operations work equally well with multidimensional arrays. Again, the arrays to be combined must have the same shape. The expressions `a+b` and `a*b` generate error messages unless

³ The module we are using is important: The `sqrt` function in the `math` module will not accept an array as an argument.

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`a.shape==b.shape` is `True` or `a` and `b` can be “broadcast” to a common shape. (For example, you can add a number to an array of any shape. For more information, try a web search for `numpy broadcasting`.)

Most math operations on NumPy arrays are performed item by item.

Vectorized operations apply only to NumPy arrays. Most mathematical operators are not defined for Python lists, tuples, or strings. Some are, but they do not carry out arithmetic. Execute the following commands to see why we usually use arrays instead of Python’s `list` data structure in numerical work:

```
x = [1, 2, 3, 4, 5, 6]
2 * x
x + 2
x * x
5 x - x
```

If you “add” two lists, two tuples, or two strings with a `+`, the result will be to join, or concatenate, them. Python’s `list` objects are useful in many computing applications, but fast mathematical calculation on large collections of numbers is not one of them.

T2 Vectorizing logic

Python’s logical operators like `and`, `or`, and `not` will give errors when applied to arrays. If you need vectorized logical operations on *Boolean* arrays, you can use arithmetic operators instead. “Addition” of `bool` objects means the logical OR, and “multiplication” means logical AND. A `~` (tilde) in front of a Boolean array means NOT, or logical negation. Try:

```
x = np.array([True, True, False, False])
y = np.array([True, False, True, False])

x + y      # apply OR element by element
5 x * y    # apply AND element by element
~x         # apply NOT element by element
```

3.2.2 Matrix math

Sometimes, instead of item-by-item operations, you may want to combine arrays by the rules of matrix math. For example, the “dot product” of two vectors (or, more generally, matrix multiplication) requires a special function call. Compare the output of the two operations below:

```
a = np.array( [1, 2, 3] )
b = np.array( [1, 0.1, 0.01] )
a*b
np.dot(a, b)
```

The number of elements in `a` equals the number of elements in `b`, so the dot product is defined. In this case, it is the single number⁴ $\mathbf{a} \cdot \mathbf{b} = a_1b_1 + a_2b_2 + a_3b_3$.

⁴ There are alternative ways to evaluate dot products. Each array has a `dot` method: `a.dot(b)` is equivalent to `np.dot(a,b)`. In Python 3.5 and later, you can also use the “at” sign as shorthand for matrix multiplication: `a@b` is also equivalent to `np.dot(a,b)`.

Your
Turn
3B

Define `a` as a row vector and `b` as a column vector by replacing the first two lines above:

```
a = np.array( [1, 2, 3] ).reshape((3,1))
b = np.array( [1, 0.1, 0.01] ).reshape((1,3))
```

Again evaluate `a*b` and `np.dot(a,b)` and explain what you get. The first result may surprise you. It is a useful operation called an “outer product.”

3.2.3 Reducing an array

In contrast to ordinary functions like `sin`, which act item by item when given an array, some functions combine elements of an array to produce a result that has fewer elements than the original array. Sometimes the result is a single number. Such operations are said to **reduce** an array.

A common array reduction is finding the sum of the elements in each row or column, or the sum of all the elements in the array. Try

```
a = np.vstack( (np.arange(20), np.arange(100, 120)) )
b = np.sum(a, 0)
c = np.sum(a, 1)
d = np.sum(a)
```

The function `np.sum(a,n)` takes an array `a` and an integer `n` and creates a new array. Each entry in the new array is the sum of the entries in `a` over all allowed values of index `n`, holding the other indices fixed. In the example above, `n=0` specifies the first axis; thus, `b` contains the column sums of `a`. Setting `n=1` specifies the second axis; thus `c` contains the row sums of `a`. When no index is given, as for `d`, `np.sum` adds up *all* of the elements in the array.

Python also has a built-in `sum` function that does not work the same way. Try the example above using `sum` without the “`np.`” prefix to see what happens, then look at `help(sum)` and `help(np.sum)` to understand the difference between the two functions.

You can use experimentation and the `help` command to explore the useful related functions `np.prod`, `np.mean`, `np.std`, `np.min`, and `np.max`. Alternatively, every array comes with methods that evaluate these functions on the array’s own data. For instance, the example above could also have been written as follows:

```
a = np.vstack( (np.arange(20), np.arange(100, 120)) )
b = a.sum(0)
c = a.sum(1)
d = a.sum()
```

See if you can explain how the following code calculates 10!:

```
ten_factorial = np.arange(1, 11).prod()
```


3.3 SCRIPTS

You can do many useful things from the IPython console, which is what we have been using up to this point. However, retyping the same lines of code at the command prompt over and over is tedious, and many tasks involve code that is far more complex than the examples above. Your code will go through many versions as you get it right, and it's tiresome to keep retyping things and making new errors. You will want to work on it, take a break, return to it, close Spyder, move to a different computer, and so on. And you will want to share code with other people in a “pure” form, free from all the typos, missteps, and output made along the way. For all these reasons, you will probably do a lot of your coding in scripts.

A **script** is simply a text file that contains a series of Python commands. You edit a script in a text editor, and then execute it by calling it with a single command or a mouse click.

3.3.1 The Editor

You can use Spyder’s Editor to create and edit scripts. To use the Editor, simply click on its pane or use the keyboard shortcut <Cmd-Shift-E>. If the Editor is not open, the keyboard shortcut will open it.

To create a script, open a new document from the File menu, click on the blank page icon at the upper left, use the keyboard shortcut <Cmd-N>, or type `%edit` in the IPython console.⁵

Write a script that contains any of the code fragments given earlier. As you enter text in a script, hitting <Return> only moves the cursor to the next line. No commands are executed because Python has no knowledge of your script yet. It is just a text file somewhere in your computer’s memory.

When you finish editing, you can execute your file by clicking the  button, selecting “Run” from the Run menu at the top of the window, or by using the shortcut key <F5>. Python executes the code in the Editor’s active tab and displays any output in the IPython console. Before it does that, however, it saves the code, prompting you for a file name if necessary. It is customary to end Python code file names with the extension `.py`.⁶

Type `%reset` at the command prompt before running a script.

That way, you know exactly what state Python is in when the script begins. Keep in mind that `%reset` will delete all imported modules. (See Section 1.2.1.) Thus, you will probably want to include the following lines at the top of *every* script you write:

```
import numpy as np
import matplotlib.pyplot as plt
```

You can have Spyder add these lines to every new script by default. See Appendix A.

3.3.2 Other editors

If you prefer another text editor, you can edit and save files outside of Spyder and then run them within Spyder. Give your code file the extension `.py` (not `.txt`). You can then type `%run` followed by a file name in the current directory (with or without the `.py` extension) at the command prompt. You will need

⁵ Keyboard shortcuts may not work unless you click inside the Editor window first.

⁶ When naming your script, do not use a Python reserved word (such as `for.py`), nor the name of a module you may want (such as `numpy.py`).

to import modules and define variables within the script instead of relying on what you may have already done in the IPython console.

See Appendix B for more information about plain text editors and working with Python from the command line.

3.3.3 First steps to debugging

If you make an error in your code while you are typing it in the Editor, Spyder may catch it before you even click [RUN]. Lines with potential errors are flagged by a yellow triangle with an exclamation point inside or a red circle with an “X” inside. A red circle means your script will not run; a yellow triangle means it may not run.⁷ Type the following lines in the Editor to see the two types of errors:

```
import numpy as np
s = str(3
prnt(s)
sqrt(3)
```

Spyder identifies a *syntax error* in the second line and flags it with a red circle. Move the mouse cursor over the red circle to get a brief explanation. The information provided is sometimes cryptic, but just knowing the general location of an error is quite helpful.

Fix the second line so that `s` is assigned the string representation of 3. Now, Spyder identifies more problems. (If there is a fatal error, Spyder will not bother warning you about other potential trouble. Thus, if there is more than one fatal error, Spyder will usually recognize only the first.)

A yellow triangle appears next to the first line, and red circles appear next to the third and fourth lines. The yellow triangle is only a warning: We have imported a module without using it. The red circles indicate more serious errors. The debugger has identified an undefined function called `prnt` in the third line, and one called `sqrt` in the fourth. If you run the script, Spyder will attempt to execute the code. However, when Python encounters the `prnt` command, it discovers that no such name is defined, and so it *raises an exception* called `NameError`. That is, it prints a *runtime error message* in the IPython console and aborts any further processing.

Python error messages can be long. The last few lines are the most helpful.

The last few lines of the error message will cite the line number where Python first noticed something wrong, display a portion of the script near this line, and describe the type of error encountered. The script may contain other errors, but Python quits when it hits the first one. Fix it and try again.

The most common cause of `NameError` is misspelling.

Variable names are case sensitive, so inconsistent case is a spelling mistake. For example, if you define `myVelocity=1` and later attempt to use `myvelocity`, Python treats the second instance as a totally new, undefined variable. Scanning the Variable Explorer can help you find such errors.

Another common cause of `NameError` is using the wrong name or nickname for a module.

⁷ [T]Spyder’s syntax checking is performed by the `pyflakes` module, which can be run independently of Spyder. Another popular

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This is what is happening in the fourth line of the script. Try to fix all the errors and get the script to run.

Runtime errors generate messages in the IPython console when you run the code. Most cause Python to halt immediately. A few, however, are “nonfatal.” For example, try

```
for x in np.arange(-1, 8):
    print(x, np.log(x))
```

Python finds nothing syntactically incorrect here, but NumPy recognizes a problem when it attempts to calculate the first two values. It gives these values names in this case (`nan` for “not a number” and `-inf` for “minus infinity”), but it knows that these values could cause problems later, so it issues two `RuntimeWarning` messages, and even explains what happened.⁸

Still other situations generate no message at all—you just get puzzling results. You will encounter many situations of this sort, and then you’ll need to *debug* your code.

Whole books are written about the art of debugging. When a code generates messages or output that you don’t understand, you need to look for clues. Here are some ideas:

- Read your code carefully. This is often the quickest route to insight.
- Build your code slowly. Most scripts execute a complicated task as a series of simple tasks. Make sure each step does exactly what it is supposed to. If every step does exactly what it is supposed to do and your code still does not work, you have a *theory bug*, not a *programming bug*.
- Start with an easier case. You’re using a computer because you can’t do the problem by hand, but maybe there’s another case that you *can* do by hand. Adapt your code for that case (perhaps just a matter of changing a few parameter values), and compare the output to the answer you know to be correct.
- Probe your variables. After a code finishes (or terminates with an error), all of its variables retain their most recent values; check them to see if anything seems amiss. (You can use the Variable Explorer or `print` statements.)
- Insert diagnostics. Somewhere prior to where you suspect there’s an error, you can add a few lines that cause Python to print out the values of some variables *at that moment* in the code’s execution. When you run the program again, you can check whether they’re what you expect.
- Be proactive as you write. Make sure that what you think *should be* true actually *is* true after a command is executed. Python offers a very useful tool for this: the `assert` statement. At any point in your code, you can insert a line of the form

```
assert (condition), "Message string"
```

If the condition is `True`, then Python will continue executing your code. If it is `False`, however, Python will stop the code, give an `AssertionError`, then print your message. For example, in the code above, we could insert the following line above the `print` statement in the most recent code sample:

```
assert (x > 0), "I do not know how to take the log of {}!".format(x)
```

When the code runs, it will not only stop when it tries to take the logarithm of a bad value—it will also print out the value that caused the problem. Of course, you can’t foresee every possible exceptional case. But you will develop a sense of what bugs are likely after some practice.

- Explain the code line-by-line out loud to another person or inanimate object. The latter practice is often called *Rubber Duck Debugging*. Professional programmers sometimes force themselves to explain

⁸ Not all modules are as forgiving. Try importing the `math` module and using its `log` function inside the loop instead.

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malfunctioning code to a rubber duck or similar totem to avoid having to involve another developer. The act of describing what the code is *supposed* to do while examining what it actually does quickly reveals discrepancies.

- Ask a more experienced friend. This may be embarrassing, because almost all coding errors appear “stupid” once you’ve found them. But it’s not as embarrassing as asking your instructor. And either one is better than endlessly banging your head against the wall. You need to become self-reliant, eventually, but it doesn’t happen all at once.
- Ask an online forum. An amazing, unexpected development in human civilization was the emergence of sites like stackoverflow.com, where people pose queries at every level, and total strangers freely help them. The turnaround time can be rather slow; however, your question may already be asked, answered, archived, and available.
- Learn about *breakpoints* and *interactive source code debuggers*. These tools go beyond the scope of this tutorial, but there may come a day when your code is complex enough to need them.

Maybe the single most important point to appreciate about debugging is that *it always takes longer than you expect*. The inevitable corollary is

Don’t wait until the day before a project is due before getting started.

Some puzzles don’t resolve until your subconscious has had time to unravel them. If you need help from a friend, lab partner, or instructor, that takes time, too. Respect the subtlety of coding, and give yourself enough time.

3.3.4 Good practice: Commenting

Another benefit of writing scripts is that you are free to include as many remarks to your reader—and yourself—as you like.

Let’s apply our example of calculating solutions to the quadratic equation by giving it some physical meaning. Suppose you are planning a prank on a friend and need to know how long a snowball remains in the air when thrown upward with a particular initial speed. You recall from introductory physics that the height of a ball thrown upward is

$$y(t) = y_0 + v_0 t - \frac{1}{2} g t^2.$$

After you recall the meaning of all the symbols, you realize that you need the value of t for which $y(t) = 0$. Rearranging things a bit, you obtain

$$\frac{1}{2} g t^2 - v_0 t - y_0 = 0.$$

This is the quadratic equation we already solved, but now the parameters all have physical meanings.

You can use the following code to plan your prank:

```
# projectile.py      [get code]
# Jesse M. Kinder -- 2021
"""
Calculate how long an object is in the air when thrown from a specified height
5 with a range of initial speeds assuming constant acceleration due to gravity:
```



```

0.5 * g * t**2 - v0 * t - y0 = 0
"""

import numpy as np
10
    %% Initialize variables.
initial_speed = 0.0          # v0 = initial vertical speed of ball [m/s]
impact_time = 0.0            # t = time of impact [s] (computed in loop)

15 %% Initialize parameters.
g = 9.80665                  # Gravitational acceleration [m/s^2]
initial_height = 2.0          # y0 = height ball is thrown from [m]
speed_increment = 5.0         # Speed increment for each iteration [m/s]
cutoff_time = 10.0            # Stop computing after impact time exceeds cutoff.

20 %% Calculate and display impact time. Increment initial speed each step.
# Repeat until impact time exceeds cutoff.
while impact_time < cutoff_time:
    # Use quadratic equation to solve kinematic equation for impact time:
    impact_time = (np.sqrt(initial_speed**2 + 2 * g * initial_height) \
                   + initial_speed) / g
    print("speed= {} m/s; time= {:.1f} s".format(initial_speed, impact_time))
    initial_speed += speed_increment
print("Calculation complete.")

```

(The syntax `x+=1` is a concise and descriptive shorthand for `x=x+1`. Other arithmetic operations can be called in a similar way. For example, `x*=2` will double the value of `x`.)

Compare this code to what was written in Section 3.1.1. The code here accomplishes a similar task, but the purpose and function of the script are now easy to understand because of the comments, whitespace, and meaningful variable names.

- The opening lines now tell us who wrote the program, why, and when.
- Next we import NumPy.
- Lines 11–19 assign values and meaningful names to the parameters and variables. (By “parameter” we simply mean a variable whose value is not going to change throughout the execution of the code.) Comments describe the role of each.
- The first parts of lines 12–13 and lines 16–19 are code, but everything after the hash signs (#) is commentary explaining the meaning of the assignments.
- Lines 21–22 introduce the main loop of the program and explain the condition for exiting the loop.

The script is certainly longer than the bare code in Section 3.1.1, but which is easier to read? Which would you rather read, if you wanted to reuse this code after a month or two of not working on it?

*Every coder eventually learns that good comments save time in the long run ...
as long as comments are updated to reflect changes in the code.*

You will probably ignore this advice until you return to a script you wrote several weeks earlier, cannot figure out what it means, and have to rewrite the whole thing from scratch. At least you were warned.

The script makes use of two different types of comments: One type is enclosed within triple quotes ("'''"), and the other is preceded by the hash sign (#). The hash sign is an *inline* comment character. Python will ignore everything that follows a hash sign on a single line. Comments can start at the beginning of a line or in the middle. Placing a short comment on the same line as the code being discussed is a good way to ensure that you'll remember to update the comment if you change the line later.

Some comments begin with the hash sign followed by two percent signs (#%). These mean nothing special to Python, but they divide the code into logical units called **cells**. Besides looking good on the screen (try it), this structuring allows you to run individual cells separately within Spyder. To see how this works, click anywhere inside such a cell, then click the **RUN CELL** button (immediately to the right of the **RUN ▶** button) or use the shortcut <Ctrl-Return>. Also try the alternative **RUN CELL AND ADVANCE** (immediately to the right of the **RUN CELL** button), which has the shortcut <Shift-Return>. Breaking code into cells by using comments is another useful debugging tool.

Triple quotes provide a convenient way to create strings that span multiple lines. They can also be used to create special kinds of comments called **documentation strings**, or **docstrings** for short. A string enclosed in triple quotes becomes a docstring when it is placed at the beginning of a file or immediately after a function declaration.⁹ Anywhere else in the file, a string enclosed in triple quotes is treated as an ordinary string.

Python uses docstrings to provide information about modules and functions you write. For example, the docstring in a function can tell a user what arguments are required, what the function will return, and so on. The script above is not a very useful module, but to see how the **help** feature works with the docstrings you write, you can import it anyway. Type **import projectile** (the file name without the .py extension) at the command prompt. Now type **help(projectile)**. You should see the text between the triple quotes.

A final comment about comments: As you hack together a workable code, you may want to try out some temporary change, but preserve the option to revert your code to its preceding version. One method is simply to *keep* the old lines of code, but “comment them out” so that they become invisible to Python. If you want them back later, simply “uncomment” them. Spyder, like most other editors, offers a menu item to comment or uncomment a selected region of text. You can also use the shortcut <Cmd-1>.¹⁰

The sample script above also illustrates proper use of indentation and other forms of *whitespace* (spaces, tabs, and blank lines).

- Proper indentation is crucial in Python. It also makes the code easier to read and understand. There is no ambiguity about which lines of code belong to the **while** loop and which do not.
- Python ignores blank lines, but they make the code much easier to read and interpret by dividing it into logical units.

3.3.5 Good practice: Using named parameters

Why did we clutter the code above with a variable name like `initial_height` instead of `y0`? Why did we bother to name the fixed parameters at all? We could have “hard coded” everything, replacing `g`, `initial_height`, and `speed_increment` by the numerical values 9.8066, 2.0, and 5.0 everywhere. There are at least four reasons:

⁹ Section 6.1 discusses user-defined functions.

¹⁰ You can also use a version control system (like Git, described in Appendix B) to clone your code, try out options, and merge the best ones back into a master version.

1. We have separated what the code does from its input. This type of abstraction allows you to easily adapt your code for other purposes. You can run the same calculation with different values for the fixed parameters, embed the main loop of this code *inside* a loop in another program that evaluates *many* values of `initial_height`, or use the main loop inside a *function* that accepts the parameter values as input. Keeping them symbolic helps.
2. Using meaningful names for variables and fixed quantities makes your code easier to read and understand. Would you be as likely to understand the purpose of the program if there were no comments and we had just used short, uninformative variable names like `x1`, `x2`, and `x3`? You can probably recall taking notes in a physics or math class and momentarily forgetting what quantity a particular symbol stands for. It is much faster to write `y0` on the chalkboard or in your notebook, but the meaning is not always clear. There is no ambiguity about what the variable `initial_height` represents.
3. Naming parameters also keeps their values distinct, even when they are numerically equal. What if there were two parameters with different physical meanings that had the same numerical value of `5.0`? If you give variables and parameters different, meaningful names, then you won't get confused by such coincidences.
4. Using named parameters to control the *size* of a calculation is extremely useful. For instance, you may intend to set up ten arrays with five million entries each, and perform a thousand calculations on the collection of arrays. Such a code could take a long time to execute. It would waste a lot of time to wait for it to execute every time you fix a small bug or add a feature. It's better to set up some parameters (perhaps called `num_elements` and `max_iterations`) near the beginning of the script that control how big the calculation is going to be. You can choose small values for the development phase, and switch to the larger desired values only when you near the final version. Moreover, you can be certain that every array in the calculation will have the same number of elements.

These points are part of a bigger theme of coding practice:

Don't duplicate. Define once, and reuse often.

That is, if the same parameter appears twice, give it a name and set its value *once*. (This idea is sometimes expressed as “Don’t repeat yourself,” or DRY for short.) If you don’t implement this principle, then when you wish to change a value, inevitably you will find and change all but one instance. The one you missed will cause problems, and it could be hard to find. Worse yet, you may accidentally change something else.

Later we will see how “don’t duplicate” can be applied not just to parameter values but also to code itself when we discuss functions in Section 6.1.

3.3.6 Good practice: Units

Most physical quantities carry units, for example, 3 cm. Python doesn’t know about units; all its values are pure numbers. If you are trying to code a problem involving a quantity *L* with dimensions of length, you’ll need to represent it by a variable `length` whose value equals *L divided by some unit*. That’s fine, as long as you are consistent everywhere about what unit to use.

You can make things easier on yourself if you include a block of comments near the beginning of your code declaring (to yourself and others) the variables *and* the units you are going to use.


```

# Variables:
#
# -----
# length      = length of the microtubule [um]
# velocity    = velocity of motor          [um/s]
5 # rate_constant = rate constant        [1/s]
...

```

(The notation um is an easy-to-type substitute for μm .) As you work on the code, you can refer back to these comments to keep yourself consistent. Such fussy hygiene will save you a lot of confusion someday.

Later on, when you start to work with data files, you need to learn from whoever created the file what units are being used to express quantities. Ideally this will be spelled out in a text file (perhaps called README.txt) accompanying the data file, or in the opening lines of the data file. (You should document data files that you create in this way.)

3.4 CONTINGENT BEHAVIOR: BRANCHING

We have now explored several ways to automate repetitive calculations. Two of these methods, `for` loops and `while` loops, used control structures of the Python language to repeat a block of code. The `while` loop above modified its behavior on the fly when it decided whether to execute its block of code again depending on the result of an intermediate calculation. Every programming language, including Python, has a more general mechanism for specifying contingent behavior, called **branching**. Try entering this code in the Editor and running it:¹¹

```

# branching.py  [get code]
""" This script illustrates branching. """

import numpy as np
5
maxTrials = 5

for trial in range(1,maxTrials+1):
    userInput = input('Pick a number: ')
10    number = float(userInput)
    if number < 0:
        print('The square root is not real.')
    else:
        print('The square root of {} is {:.4f}.'.format(number,np.sqrt(number)))
15    userAgain = input('Try another [y/n]? ')
    if userAgain != 'y':
        break

    %% Check to see if the loop exited normally.
20    if trial >= maxTrials:
        print('Sorry, only {} per customer.'.format(maxTrials))

```

¹¹ In Python 2, `input` attempts to evaluate its argument as a Python expression. To run this script, users of Python 2 should replace `input` with `raw_input` or redefine `input` as `raw_input`. See Appendix E.


```

25 elif userAgain == 'n':
    print('Bye!')
else:
    print('Sorry, I did not understand that.')

```

3.4.1 The if statement

The code above illustrates several new ideas:

- The `input` function displays a prompt in the IPython console, then waits for you to type something followed by <Return>. Its return value is a string containing that input.¹² In this case, we assign that string to a variable called `userInput`, then convert it to a `float` for use later in the program.
- Line 11 contains an **if statement**. The keyword `if` is followed by a logical (*Boolean*) expression and a colon. Python evaluates this expression. If it is `True`, Python executes the next indented block of code (in this case, a single line). If the expression is `False`, Python skips the block of code.
- The conditional block of the first `if` statement is followed by the keyword `else` and a colon. If the condition in the `if` statement is `True`, then Python skips the indented block following the `else` statement on line 13. If the condition is `False`, then Python executes the indented block following the `else` statement.
- The `input` function appears again in line 15. This time, the string returned by the function is what we want, so no conversion is necessary.
- The notation `!=` in line 16 means “is not equal to.” You can also use the keyword `not` to negate a Boolean value, so we could have written `if not(userAgain=='y')`.
- Sometimes a loop should terminate prematurely (before the condition in its `for` or `while` statement is satisfied). In line 17, the keyword `break` instructs Python to exit the indented block of the `for` loop and proceed from line 18, regardless of whether the loop is done.
- Line 20 starts a three-way branch whose behavior depends on how the loop terminated. This is accomplished using the `elif` statement, short for “`else if`.” There can be as many `elif` statements as you like. Python will drop down from the opening `if` statement through the `elif` statements until it finds one that evaluates to `True` or it reaches an `else` statement. In any circumstance, only one of lines 21, 23, or 25 will be executed.

In short,

A branch construction starts with an `if` statement, may continue with a series of `elif` statements, and may conclude with an `else` statement. Only the `if` statement is required.

Each `if`, `elif`, and `else` statement is followed by an indented block of code. At most, one of these blocks will be executed. Include an `else` statement when you want *something* to happen, even when none of the `if` or `elif` statements are satisfied.

The code listed above used the relation operators `<`, `==`, and `!=` to generate Boolean (`True` or `False`) values. Other operators that return Boolean values include `>`, `<=`, and `>=`. You can generate more complex conditionals with the Boolean operators `and`, `or`, and `not`.

¹² See footnote 11.

Conditional statements with arrays

Sometimes, you may wish to use arrays in conditional statements. However, `if`, `elif`, and `else` statements require statements that evaluate to a single Boolean value: `True` or `False`. Two NumPy functions are helpful: `np.any` and `np.all`.

Suppose you have an array of numbers, `x`, and you wish to work with their logarithms. As we saw earlier, `np.log(x)` will not raise any exceptions, even if `x` contains zeros or negative values. Instead, it will return an array with `nan` and `inf` elements that might ruin subsequent calculations. We can test for this with an `if` statement in two ways. We can check to see if *any* of the values are nonpositive:

```
if np.any(x<=0):
    print("This array is dangerous for logarithms.")
else:
    print("This array is safe for logarithms.")
```

Alternatively, we can check to make sure that *all* of the values are positive:

```
if np.all(x>0):
    print("This array is safe for logarithms.")
else:
    print("This array is dangerous for logarithms.")
```

Every array has its own `any` and `all` methods. However, in Python, any number other than 0 will evaluate to `True`. Thus, `x.any()` is equivalent to `np.any(x!=0)`—*not* `np.any(x<=0)`.

3.4.2 Testing equality of floats

Numerical comparisons involving floating point numbers can be tricky. You are unlikely to ask Python whether two floats representing physical quantities are exactly equal via `a==b`. (Physical effects such as thermal motion, quantum fluctuation, and chaos ensure that this is never the case.) However, consider the following loop:

```
m, k = 0.0, 0.3
while m != k:
    print(m)
    m = m + 0.1
```

It looks harmless enough, but this loop never terminates! On the third iteration, the computer's internal representation of `m` is almost, but not quite, the same as its representation of `k`.¹³ Python continues printing indefinitely because `m` is *always* not equal to `k`.

The cure for this particular loop is to use an inequality: `while m<=k: ...`. However, you may run into a related problem if you test for equality, thinking that two variables are integers when, in fact, one or both have been converted to floats.

Never compare floats with == or !=. Compare integers, or use inequalities.

If you need to compare two floats, you should probably compare their difference to within some reasonable tolerance appropriate for the problem: `if abs(a-b) < 1e-6: ...`. NumPy also provides a

¹³  The binary representation of 1/10 is periodic: 0.0001100110011.... This cannot be accurately represented with any finite number of bits, and Python only uses 64. Thus, due to roundoff error, `0.0+0.1+0.1+0.1` will not evaluate to exactly the same floating point number as `0.3`.

convenient function to test for “reasonable agreement” between floats, and even allows you to define what is reasonable: See `help(np.isclose)`.

On truth

Python defines `True` rather broadly. Any nonzero numerical value evaluates to `True` upon conversion to type `bool`, as does any nonempty list, string, tuple, or array. The following expressions evaluate to `False`: `False`, `None`, `[]`, `()`, `{}`, `0`, `0.0`, `0j`, `" "`, `' '`. Almost everything else evaluates to `True`.

3.5 NESTING

In many cases, we may wish to embed one `for` loop inside another `for` loop, embed a `for` loop within a `while` loop, or create some other combination. For example, when dealing with probability, we sometimes wish to create an array `A` where the value of `A[m, n]` depends on the indices `m` and `n`. (`m` and `n` might be the number of times two different kinds of random event—with probabilities `p` and `q`, respectively—occur during an experiment.) We can do this with a code like the following:

```
# nesting.py      [get code]
rows = 3
columns = 4
p = 0.1
5 q = 0.3
A = np.zeros( (rows, columns) )
for m in range(rows):
    for n in range(columns):
        A[m, n] = p**m * q**n
```

Notice the following:

- Before the loops, the code creates an array with `np.zeros`. NumPy does not build arrays on the fly. It sets aside a block in the computer’s memory to store the array, so it needs to know how big that block should be.¹⁴ Thus, we first create an array with the right shape, and then assign it the values we need.
- Python’s `range` function is used to iterate over values of `m` and `n`. It is similar to `np.arange`, but, instead of an array, it creates an object that generates values as they are needed.¹⁵

Use `np.arange` when you need an array. Use `range` in `for` loops.

- Line 9 sits inside *two for* loops, so it is executed $3 \times 4 = 12$ times. One loop inside another is called a **nested loop**. Note that `if` statements and `while` loops can also be nested.
- Line 9 must be executed inside the inner loop, so it is indented twice.
- Descriptive variable names and Python’s forced use of indentation make comments nearly superfluous in this code fragment. A reader might only wonder why you are computing $A_{m+1,n+1} = p^m \cdot q^n$ and not some other function.

¹⁴ You can later resize the array if you need to, but this usually indicates poor planning. If you need a data structure with a variable size, you might use a Python list instead.

¹⁵  The `range` function in Python 3 is equivalent to the `xrange` function in Python 2. In Python 2, `range` creates an actual list of integers. It will work in `for` loops, but `xrange` is often preferable, especially for large loops. There is no function called `xrange` in Python 3.

CHAPTER 4

Data In, Results Out

Computers in the future may ... perhaps only weigh 1.5 tons.

— Popular Mechanics magazine, 1949

Most data sets are too big to enter by hand, often because they were generated by automated instruments. You need to know how to bring such a data set into your Python computing session (*import* it). You will also want to save your own work to files (*export* it) so that you do not have to repeat complex calculations. Python offers simple and efficient tools for reading and writing files.

Also, most results are too complex to grasp if presented as tables of numbers. You need to know how to present results in a graphical form that humans (including you) can understand. The PyPlot module provides an extensive collection of resources for visualizing data sets.¹

In this chapter, you will learn to

- Load data from a file;
- Save data to a file; and
- Create plots from a data set.

4.1 IMPORTING DATA

Much scientific work involves collections of experimental data, or *data sets*. In order to crunch a data set, Python must first import it. Many data sets are available in plain text files, including

- *Comma-separated value (.csv)* files: Each line of the file represents a row of an array, with entries in that row separated by commas.
- *Tab-separated value (.tsv)* files: Each line represents a row of an array, with entries in that row separated by tabs, spaces, or some combination of the two. Spacing does not have to be consistent. Python treats any amount of whitespace between entries as a single separator.

The file extensions `.txt` and `.dat` are also used for data files with comma or whitespace separation. Python has built-in support for reading and writing such files, and NumPy includes additional tools for loading and saving array data to files.

The SciPy library provides a module called `scipy.io` that will allow you to read and write data in a variety of formats, including MATLAB files, IDL files, and `.wav` sound files. See docs.scipy.org.

¹ [T] All of the operations described in this chapter—and much more—can be accomplished with the `pandas` module (see Chapter 10).

4.1.1 Obtaining data

Before you can import a data set, you must obtain the file of interest and place it in a location where Python can find it. It is good practice to keep all of the files associated with a project—data, scripts, supporting information, and the reports you produce—in a single folder (also called a **directory**).

You can tell Spyder to use a particular folder by setting the **current working directory** in the Preferences menu.² Select **Current working directory** from the choices on the left. On the right select “**the following directory:**”. Then choose a folder³ and click **APPLY** then **OK**. Close Spyder and relaunch. Now each time you launch Spyder, it will start in this folder. This is also where Spyder will attempt to load and save all data files by default. The IPython command `%run myfile.py` will also look in this folder.

It is possible to switch to a different working directory during a session in Spyder. We recommend keeping things simple by working in a single folder.

Once you have set up your working directory, you can place data files into it. To download the data sets described in this tutorial, go to

press.princeton.edu/titles/32489.html

Here, you will find a link to a zipped file that contains a collection of data sets.⁴ Follow the instructions to save the zipped file to your computer. It will probably be saved into your **Downloads** folder unless you specify a different destination. Double-click on the file to unzip it. This will create a new folder called **PMLSdata** in the current folder. You will find all of the data sets described in this tutorial, plus several more. Move **PMLSdata** to a permanent location. If you place **PMLSdata** in your working directory, you will be able to access it easily from within Spyder.

You do not have to limit yourself to the data sets described here. Someday, you’ll use data from another source. You may have an instrument in the lab that creates it, or you may get it from a public repository or a coworker. Scientific publications that contain quantitative data in the form of graphs are another source. A graph is a visual representation of a set of numbers, and can be converted back to numbers with a special purpose application. Applications that convert graphs to numbers include

- Engauge Digitizer (markummitchell.github.io/engauge-digitizer/),
- Plot Digitizer (plotdigitizer.sourceforge.net),
- DataThief (www.softpedia.com/get/Science-CAD/DataThief.shtml), and
- GetData (www.getdata-graph-digitizer.com).

4.1.2 Bringing data into Python

After you have obtained the data sets, find the entry called **01HIVseries**. This folder contains a file called **HIVseries.csv**. There is also an information file called **README.txt** that describes the data set. Copy these two files into your working directory.

Arrays

You are now ready to launch Python and load the data set. First, inspect the data using the Editor by

² One way to access the Preferences is by clicking the wrench icon (Figure 1.1, page 7).

³ We recommend a generic name, like **scratch** or **curr**. After you are done with a project, you can archive the files under a different name and create a new scratch folder.

⁴ Resources mentioned here are also maintained at a mirror site: physicalmodelingwithpython.blogspot.com. Readers familiar with Git can also obtain the files from our repository on GitHub. See Appendix B.

typing `%edit HIVseries.csv` at the IPython command line. (You can also open files in the Editor by using the `File>Open` menu option or by clicking in the Editor window and using `<Cmd-O>`, but you may need to tell Spyder to look for “all files” instead of just Python files.) You should see 16 lines of data, each containing two numbers separated by a comma. You can open the information file `README.txt` to find out what the numbers mean.

To load the data into a NumPy array, you can use the `np.loadtxt` command. This command attempts to read a text file and transform the data into an array. By default, NumPy assumes that data entries are separated by spaces and/or tabs. To load a `.csv` file, you must instruct `np.loadtxt` to use a comma as the delimiter. A look at the help file (try typing `np.loadtxt?`) shows that you can do this by including a *keyword argument* called `delimiter`. Keyword arguments are optional arguments you can pass to a function to modify its behavior (see Section 1.4.4).⁵

Thus, to load the data, use the command

```
data_set = np.loadtxt("HIVseries.csv", delimiter=',')
```

Notice how this command is used:

1. The function returns an array. We assign the variable name `data_set` to this array.
2. We give `np.loadtxt` the file name as a string.⁶
3. We specify the delimiter as a string, in this case `', '`.

Python will look for the file in the current working directory. In Spyder, you can see the working directory in a text box in the upper-right corner of the screen. You can use the IPython magic command `%pwd` to display the present working directory’s name to the screen. If the file fails to load, either it is located in the wrong folder or Spyder is looking in the wrong folder. If Spyder is no longer in its default working directory, you may need to restart Spyder and try again. If the file is in the wrong location, move it to the working directory and try to load it again.

An alternative to moving the file is to specify its complete `path`. For example, on a Mac where the `PMLSdata` folder is still in the `Downloads` folder, you could write⁷

```
data_file = "/Users/username/Downloads/PMLSdata/01HIVseries/HIVseries.csv"
data_set = np.loadtxt(data_file, delimiter=',')
```

Path names are formatted differently in Windows:

```
data_file = r"c:\windows\Downloads\PMLSdata\01HIVseries\HIVseries.csv"
```

Note the use of a raw string to handle the backslashes. (See Section 2.3.1.)

If you are working with multiple files outside the current working directory, specifying the complete path can lead to a lot of typing. Also, you might move the data to a different folder at some point in the future. For these reasons, it is often convenient to “define once, reuse often” in scripts. For example, with the following two commands near the top of the script, you can easily load multiple data files from the same folder:

⁵ By default, `np.loadtxt` will ignore any line in the file that starts with `#`. You can adjust which header lines are discarded by changing the default values of the keyword arguments `comments` and `skiprows`.

⁶ You can provide a Python file object instead, such as one created by Python’s `open` command, or the `urlopen` command in the `urllib.request` module for remotely hosted data.

⁷ Substitute your own username in the place indicated. If you do not know a file’s full path, find the file in Finder or File Explorer

and drag it into the IPython console. You can then copy and paste the file's path into your script.

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```
home_dir = "/Users/username/"
data_dir = home_dir + "Downloads/PMLSdata/01HIVseries/"
data_set = np.loadtxt(data_dir + "HIVseries.csv", delimiter=',')
```

Later, if you work on a different computer (where `home_dir` will change) or move the data on your own computer (where `data_dir` will change), you will only have to modify one or two lines to get your code running again.

Once you have successfully imported the data, inspect the array. It has two columns. You can see its size and its entries in the Variable Explorer, or by using the IPython console. There is no description of what the numbers mean. For this, you need to consult `README.txt`. (When you create your own data sets, include documentation and descriptive comments.)

T2 Other kinds of text

The data file you need to load may not be in any conveniently delimited format. One way to handle such files is to process the file line by line. The following will generate the same array as `np.loadtxt`, but it can be adapted to more exotic formats.

```
# import_text.py      [get code]
my_file = open("HIVseries.csv")
temp_data = []
for line in my_file:
    print(line)
    x, y = line.split(',')
    temp_data += [ (float(x), float(y)) ]
my_file.close()
data_set = np.array(temp_data)
```

The second line creates an object that can read the data file.⁸ The third line creates an empty list to store the data. (We use a list instead of an array because we do not know how many lines the file contains.) The `for` loop uses a convenient Python construction to process the file one line at a time. `line` is a string that contains the current line of the file. It is updated during each iteration of the loop. Inside the loop, Python displays the line being processed, and then uses the string method `line.split(',')` to break the line into multiple strings using the comma as a delimiter. Next, it converts each individual string to a number and stores the data from each line as a new ordered pair at the end of the list of data points. The loop will terminate once it reaches the end of the file. After the loop exits, lines 8–9 close the file and convert the list of data points to a two-dimensional NumPy array.

The block of code within the `for` loop that processes the data file can be modified to extract data from other types of text files.

T2 Direct import from the web

Instead of saving a data set to your computer and then reading that file, you can combine these steps and read a file directly from the web. Try the following:

```
from urllib.request import urlopen
web_file = urlopen( "https://www.physics.upenn.edu/biophys/" + \
                    "PMLS/Datasets/01HIVseries/HIVseries.csv" )
data_set = np.loadtxt(web_file, delimiter=',')
```

⁸ There is no `file` object type in Python 3, but `open('temp.txt')` returns an object similar to a Python 2 `file` object.

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After the call to `urlopen`, Python can read the data in `web_file` by using `np.loadtxt` (or any other method for processing text files), as if the data set were located in a local file.⁹

MATLAB data files

You can import data from and export data to MATLAB formats using the `scipy.io` module. The `loadmat` function will load a `.m` file into a Python dictionary whose keys are the variable names and whose values are the associated matrices. The related function `savemat` function will write a Python dictionary of variable names and associated arrays to a `.m` file. (Dictionaries are described in Section 10.1.1.)

There is an important caveat: `scipy.io` does not support the HDF5 file format introduced in MATLAB 7.3. If you try to import a file of this type using `loadmat`, you will get an error. The simplest solution, if you have access to MATLAB, is to save the file to a format supported by `scipy.io`. In MATLAB, type `save('data.mat', '-v7')`. You should be able to load `data.mat` into Python with `loadmat`.

If you do not have access to MATLAB, you can try the `mat73` package: github.com/skjerns/mat73. It is not part of the Anaconda distribution. First use conda to install the pip Python package manager from the command line: `conda install pip`. Then use pip to install `mat73` from the command line: `pip install mat73`. This module provides a `loadmat` function whose interface and behavior mirrors `scipy.io.loadmat`, but works with the new MATLAB file format. If you need to save to a MATLAB format, use the `savemat` function from `scipy.io`.

4.2 EXPORTING DATA

Saving your work is always important when working on a computer. It is even more important when you are working in the IPython console.

Data on the screen is fleeting. Data saved in files is permanent.

If you have been working for hours and finally finish crunching numbers and making plots, great! But the moment you quit Spyder, all that work is gone. Only data and code that you have saved to files will outlast your session.¹⁰

4.2.1 Scripts

Scripts are a good way to save your work. (See Section 3.3.) If you keep adding lines to a script as you work through a problem, then you will have a record of your work *and* a program that will reconstruct the state of the system right up to the last line of the script. It can build and fill arrays, load data, carry out complicated analyses, create plots, and much more. (If you have been working at the IPython command line, copying and pasting commands from Spyder’s “History log” can help turn your interactive session into a reusable script.)

Each time you run a script, Spyder first saves it. Unless you have modified the current working directory in the Preferences, however, the default folder is probably not what you want. You can use `File>Save As...` to specify the destination. Be sure you know where your script is saved. (See Section 4.1.1 and Appendix A.)

⁹  In Python 2, the relevant command is `from urllib import urlopen`.

¹⁰ You may be able to use IPython’s history to reconstruct your work. See Section 3.1.4 (page 37).

You can also store all or part of your script in other files. Because scripts are plain text files, you can easily copy and paste code into a word processor, a homework assignment, your personal coding log, or an e-mail.

4.2.2 Data files

Python does *not* automatically save its state or the values of any variables created during a session. Hence, the importance of writing scripts: They can reconstruct the state from scratch. However, with large data sets and complex analyses, rerunning the script could take a long time. Once you have generated the data you need, you can save it to a file and simply load the file the next time you want to use the data.

Data to be read by Python

NumPy provides three convenient functions for saving data stored in arrays to a file:

- **`np.save`** – save a single array as a NumPy archive file (not human readable), with extension `.npy`
- **`np.savez`** – save multiple arrays as a single NumPy archive file, with extension `.npz`
- **`np.savetxt`** – save a single array as a text file (human readable), with an extension of your choice.

For example, the following code will store two arrays in five different files:

```
# save_load.py      [get code]
x = np.linspace(0, 1, 1001)
y = 3*np.sin(x)**3 - np.sin(x)

5 np.save('x_values', x)
np.save('y_values', y)
np.savetxt('x_values.dat', x)
np.savetxt('y_values.dat', y)
np.savez('xy_values', x_vals=x, y_vals=y)
```

Explore the resulting files, which have been created in the current working directory. Those created by `np.save` and `np.savez` have the extensions `.npy` and `.npz`, respectively, whereas those created by `np.savetxt` have the `.dat` extension we provided. (They contain tab-separated values. To generate comma-separated values, again you must use the `delimiter` keyword argument.)

You can view any of these files by opening them in a text editor. Do not save any changes to a `.npy` or `.npz` file when you close it—that will corrupt the data. If you need to show your results to someone else, or use them outside of Python, use `np.savetxt`. As you will see, `.npy` and `.npz` files are not easy to read.

To recover the saved data at any point—in the current session or in the future—use `np.load` or `np.loadtxt`, depending on the file type:

```
x2 = np.load('x_values.npy')
y2 = np.loadtxt('y_values.dat')
w = np.load('xy_values.npz')
```

The variables `x2` and `y2` now refer to arrays containing the same data as `x` and `y`. In the third example, `w` is not an array at all; however, our data is stored inside it.¹¹ If you type `w.files`, Python will return a list with two elements: the two strings '`x_vals`' and '`y_vals`'. These are the names we provided as

¹¹  It is a special object whose contents can be accessed like those of a Python **dictionary**. See Section 10.1.1.

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keyword arguments in the strange-looking call to `np.savetxt` above. We can access our data from `w` by using these keys. The following commands show that the data have been faithfully saved and loaded:

```
x2 == x
y2 == y
w['x_vals'] == x
w['y_vals'] == y
```

From these examples, you can see that `np.savetxt` offers a convenient way to save several arrays in a single file.¹² *Be sure you provide descriptive names for your arrays as keyword arguments.* If you do not, NumPy will use the unenlightening names “`arr_0`, `arr_1`, ...”

Data to be read by humans

Not all the data you generate will be in arrays. Even with array data, sometimes you will need to write information to a text file, and you may want more control over its format than `np.savetxt` offers. (Perhaps the file will be read by a colleague, or by an application other than Python.)

Recall Section 3.3.4, where snowball impact times were displayed in the console. When the session ends, this information will be lost unless you write it to a file. For small amounts of information, you can simply copy from the IPython console (or History log) and paste into a text editor or word processor.

For large data sets, you will want to write data to a file *without* displaying it on the screen. Python’s built-in function `open` will allow you to write directly to a file. To illustrate the method, let’s display the first ten powers of 2 and 3 *and* store these data for later use. The following script accomplishes both tasks:

```
# print_write.py      [get code]
my_file = open('power.txt', 'w')
print( " N \t\ t2**N\t\ t3**N" )           # Print labels for columns.
print( "----\t\ t----\t\ t----" )           # Print separator.
5 my_file.write( " N \t\ t2**N\t\ t3**N\n" )# Write labels to file.
my_file.write( "----\t\ t----\t\ t----\n" )# Write separator to file.
#%% Loop over integers from 0 to 10 and print/write results.
for N in range(11):
    print( "{:d}\t\ t{:d}\t\ t{:d}" .format(N, pow(2,N), pow(3,N)) )
10   my_file.write( "{:d}\t\ t{:d}\t\ t{:d}\n" .format(N, pow(2,N), pow(3,N)) )
my_file.close()
```

Compare the output displayed to the screen with the content of `power.txt`.

Writing text to a file is similar to printing to the display.

The differences are that

- You must open a file before writing to it: `my_file=open('file.txt','w')`. (The ‘`w`’ option means this file will be opened for writing. This erases any existing file of the same name *without* any warning or request for confirmation.)
- You must explicitly tell Python where you want a line to start and end. Wherever you have typed ‘`\n`’, Python will insert a new line. (Similarly, wherever you have typed ‘`\t`’, Python will insert a tab.)
- When you are done writing, you should close a file with the command `my_file.close()`.

¹² To reduce the file size, use `np.savez_compressed` instead.

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Python is a powerful tool for reading, writing, and manipulating text files, but we will not explore these capabilities any further. You can already accomplish quite a lot using the basic commands above in combination with the string formatting methods of Section 2.3.

What should you name your file? Python will accept any valid string you type, but your operating system may not be as flexible. It is good practice to restrict filenames to letters, numbers, underscore (`_`), hyphen (`-`), and period.

4.3 VISUALIZING DATA

With data in hand, we are ready for graphics. Python has no built-in graphing functions. Almost all of the graphing tools we discuss come from the PyPlot module, which is part of the larger Matplotlib module. To gain access to these functions, you must import the PyPlot module:

```
import matplotlib.pyplot as plt
```

4.3.1 The `plot` command and its relatives

PyPlot will make an ordinary, two-dimensional graph for you if you supply it with a set of (x, y) pairs. It's up to you to space those points appropriately. Try

```
# simple_plot.py      [get code]
import numpy as np, matplotlib.pyplot as plt
num_points = 5
x_min, x_max = 0, 4
5 x_values = np.linspace(x_min, x_max, num_points)
y_values = x_values**2
plt.plot(x_values, y_values)
# plt.show()
```

The last line may be necessary in your Python environment. If no figure window opens, try uncommenting the last line or typing `plt.show()` at the IPython command prompt.

If you are doing a lot of plotting from the IPython command line, you may wish to use the command `plt.ion()` to turn interactive plotting **on**. This will update the figure every time you execute a plotting command, but it may be slow for complex graphics. You can turn interactive plotting off by calling `plt.ioff()`.

The figure window may appear automatically, but it can still be hard to find. Try hiding your other applications and moving the Spyder window around. Your plot may be behind it. Changing to a different graphics *back end* may help.¹³ (See Section A.2.2, page 163.)

¹³ In the blog that accompanies this book, we describe a way to raise a figure window to the foreground using the Qt back end. (See press.princeton.edu/titles/32489.html.)

The code given above is a basic plotting script. Notice what happens when you execute it:

- A new window appears. (See Section A.2 if it does not.) This is now the *current figure*. Python will name it “Figure 1” if no other figure windows are open.
- Inside the current figure, PyPlot has drawn and labeled some axes. These are now the *current axes*.
- PyPlot has automatically drawn a region of the xy plane that contains all the points you supplied.
- The `plot` function takes the items in its first argument one at a time, combines them with the corresponding items in the second argument to make (x, y) pairs, and connects those points by a solid blue line. This graph is jagged, but you can fix that by specifying a larger value for `num_points`. Try it.
- With some back ends, the figure window remains “live” until you close it. That is, you can continue to modify the figure by issuing additional commands at the IPython prompt. Other back ends may force you to close a figure before you can type any more commands. See Section A.2.2 for instructions on how to change the back end.
- Changing the data in `y_values` will not automatically update your plot. You must issue another `plot` command to see such changes—or use the `set_data` method described in Section 4.3.4 below.
- You can close a figure window manually in the usual way (clicking on a red dot or “X” in the corner of the figure window), or from the IPython command prompt with the command `plt.close()`. You can close all open figure windows with the command `plt.close('all')`.

The two arrays that you supply to `plt.plot` must be of exactly the same length. That may sound obvious, but it’s surprising how often you can get just one extra element in one of them. If you’re not sure exactly how many elements you’ll get from `a=np.arange(0, 26, 0.13)`, just take a few seconds to type `len(a)` at the IPython console prompt and find out. (The function `len` is Python’s built-in function for counting the number of elements in a data structure.) You can also use `np.shape(a)`, to ask an array about its shape, or request the data field `a.shape`.

Check the lengths of your arrays.

Python’s `assert` command provides a useful test (Section 3.3.3, page 43). For the example above, you could insert a single statement before the plot command:

```
assert len(x_values) == len(y_values), \
    "Length-mismatch: {:d} versus {:d}".format(len(x_values), len(y_values))
plt.plot(x_values, y_values)
```

PyPlot will issue its own error message if you try to plot lists of unequal lengths, but an `assert` statement provides you with more information. To see the difference, rewrite the script to change the length of `x_values` after defining `y_values` and run the script with and without the `assert` statement.

You can avoid many of these length mismatch errors by adhering to the principle of “Define once, and reuse often.” For example, if the length of every array is determined by the single parameter `num_points`, then they will all have the same length.

Plotting options

Every visual aspect of a graph can be changed from its default. You can specify these when you create a

plot; most can also be changed later. Unless you only want to look at a rough plot of some data once and then discard the graph, you should write a script to generate your plot. Otherwise, you will find yourself retying a lot of commands just to change a label or the color of a single curve in the plot. Here are some of the more common adjustments you can make:

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- You can change the color of a curve when you first plot it by adding an optional argument:
`plt.plot(x_values,y_values,'r')` produces a solid red line. Other line and marker styles include
 - '`b`' (blue), '`k`' (black), '`g`' (green), and so on.
 - '`:`' (join data points with a dotted line), '`--`' (dashed line), '`-`' (solid line), and so on. Remember that red, blue, and other colors look similar on a grayscale printout. Dotted and dashed lines are much easier to distinguish.¹⁴
 - '`.`' (small dot at each data point), '`o`' (larger dot), and so on. For an open circle, use '`o`', `markerfacecolor='none'`. For other marker sizes, use the `markersize` keyword argument.

To an extent, options can be combined: For example, `plt.plot(x_values,y_values,'r--o')` plots a red dashed line with red circles at each point. Python's default is to draw a solid line and no markers. If you specify a marker style, but no line style, then no line will be drawn.

If you are plotting many lines or points on the same axes, you can use the `alpha` keyword to make them partially transparent and easier to see: `alpha=0.0` is invisible, and `alpha=1.0` is opaque.

- You can change Python's choice of plot region even *after* creating the plot. Use the command `plt.xlim(1,6)` to display the region $1 \leq x \leq 6$, using the default for y . Similarly, `plt.ylim` adjusts the vertical region. Both functions will also accept keyword arguments,¹⁵ or a list, tuple, or array containing the lowest and highest values to display on the axis: `plt.xlim(xmax=6, xmin=1)`, `plt.xlim([1,6])`, `plt.xlim((1,6))`.
- The command `plt.axis('tight')` effectively issues `xlim` and `ylim` commands to make the axes just fit the range of the data, without extra space around it. (Note the spelling: `plt.axis`, with an 'i', modifies the current axes; `plt.axes`, with an 'e', creates new axes.)
- PyPlot determines the height and width of the figure and then scales the x and y axes by different amounts to make everything fit into the figure. You may not like the distortion of your figure that arises from scaling x and y independently. The command `plt.axis('equal')` forces each axis to use the same scaling; that is, equal coordinate intervals give equal distances along each axis. Without this option, a circle may come out looking like an ellipse. The command `plt.axis('square')` does the same thing, but also adjusts the coordinate limits so that the graph is actually square in shape.

The documentation on `plt.plot` is quite helpful. Type `help(plt.plot)` or `plt.plot?` at the command prompt to read it. It describes the available marker and line styles, describes other adjustable parameters, and provides several examples.

Some of the commands just mentioned illustrate common themes: Plot properties can be set by passing strings as arguments to `plt.plot`. Some must appear in a specific position in the list of arguments, like '`r`'. Alternatively, any data field can be modified by pairing a keyword with the desired value, as in `linestyle='r--'` or its abbreviated form `ls='r--'`.

If you don't want to join your plotted points with a line, there is a separate function called `plt.scatter` that can be called instead of using the options listed above. It gives you more control over the size, style, and color of individual marker symbols than `plt.plot`.

¹⁴ Remember, too, that some readers may be colorblind.

¹⁵ Keyword arguments were introduced in Section 1.4.4 and will be discussed further in Section 6.1.3.

4.3.2 Log axes

If you'd prefer a logarithmic vertical axis, use `plt.semilogy` in place of `plt.plot`. You can also call `plt.semilogy()` after creating a plot. For a logarithmic horizontal axis, use `plt.semilogx`; for a log-log plot, use `plt.loglog`. In the two latter cases, you may want to evaluate your function at a set of values that are uniformly spaced on a logarithmic scale. See `help(np.logspace)` for a function that creates such arrays.

Your
Turn
4A

- Plot the functions $\exp(x)$ and $x^{3.6}$ over the range $2 \leq x \leq 7$. The two curves should look similar.
- Now, make a semilog plot of the same two functions. How do the two curves differ in this representation?
- Finally, make a log-log plot of the same two functions. How do the two curves differ in this representation?

You have just seen that

Linear, semilog, and log-log plots can help discriminate different relationships in data sets.

4.3.3 Manipulate and embellish

There is no limit to the time you can spend improving your graphs. Here we introduce a few more useful adjustments. You can locate many others by using your favorite search engine. (Try searching the web for “python graph triangle symbol”.) Just remember: When you close a figure, your graph is gone. However, if you write a script to construct the figure and add all of your embellishments, you can easily reconstruct it later.

The commands below may look strange at first, but you will get used to them with practice, especially if you record the ones you use in your coding log (see Section 1.2.4).

You see a plot as an image on your screen, but to Python, a plot is an *object* with many associated *attributes*: *data fields* to store plot information and properties, and *methods* for modifying those data. To adjust the properties of an existing figure, first assign a variable to the plot object:

```
# graph_modifications.py      [get code]
ax = plt.gca()
```

In this command, `plt.gca()` (get current axes) returns an object that controls many of the attributes of the plot in the current figure. It is called an *Axes* object, and it includes the axes you see, tick marks, labels, plot data like curves and symbols, and much more. An *Axes* object is fairly complex, with more than 450 properties and methods! However, you can accomplish a lot by using only a few of these. The commands fall into two general categories: *getting* data from the object and *setting* values of properties that you want to adjust. These operations have fairly descriptive names, like `ax.get_xticks()` and `ax.set_xlabel()`. Many of these operations can also be accomplished with functions from the PyPlot module itself.

Here are a few useful operations.¹⁶

¹⁶ The following commands were tested on the Qt5Agg back end. They may not work correctly with all graphics back ends.

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Title: You can add a title with `ax.set_title("My first plot")`. If you don't like the default font, you can change it by using keyword arguments:

```
ax.set_title("My first plot", size=24, weight='bold')
```

Alternatively, you may use

```
plt.title("My first plot", size=24, weight='bold')
```

Axis labels: You can—and should—label the axes with

```
ax.set_xlabel("speed")
ax.set_ylabel("kinetic energy")
```

or

```
plt.xlabel("speed")
plt.ylabel("kinetic energy")
```

Even better, you can—and should—include units: `ax.set_xlabel("speed [um/s]")`. Again, “um” is an informal abbreviation for micrometer. But it’s almost as easy to write μm instead, which will produce the standard abbreviation μm . (See Section 2.3.) If you are reporting a dimensionless quantity, like concentration relative to its value at time zero, help your readers by stating “ $c/c(0)$ [unitless]” or “concentration [a.u.]”.

Tick labels: You can change the font and size of the numbers labeling the tick marks along each axis by first creating the plot, then giving the commands

```
ax.set_xticklabels(ax.get_xticks(), family='monospace', fontsize=10)
ax.set_yticklabels(ax.get_yticks(), family='monospace', fontsize=10)
```

Line style: You can change the properties of lines after the initial plot command has been executed.

First, you need to gain access to the objects Python uses to draw the lines, then **set** their **properties** using PyPlot’s `plt.setp` command:

```
lines = ax.get_lines()      # Lines is a list of line objects.
# Make the first line thick, dashed, and red.
plt.setp(lines[0], linestyle='--', linewidth=3, color='r')
```

Legend: You can add a **legend**—a descriptive text label for each line. You can do this in two ways: Give each line a label when it is created by using the `label` keyword, or use the `set_label` method of a line object.

```
# Use "label" keyword to set labels when plotting.
plt.plot(x_values, y_values, label="Population 1")
plt.plot(x_values, x_values**3, label="Population 2")
plt.legend()                  # Display legend in plot.

# Use line objects to set labels after plotting.
ax = plt.gca()
lines = ax.get_lines()
lines[0].set_label("Infected Population")
lines[1].set_label("Cured Population")
```

```
10 ax.legend() # Display legend in plot.
```

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No legend will appear in your plot until you call `ax.legend()` or `plt.legend()`.

Text boxes: Try:

```
plt.text(2, 3, "y = {:.3f} x + {:.3f}".format(slope, intercept))
```

The first two arguments give a location, in data coordinates. The third argument is a string to be displayed at that location. (You will need to supply values for the slope and intercept.)

These examples show that you can add a lot of useful information to a plot and make it visually appealing. You can also see that this involves a lot of typing. Once you have your plot looking the way you want—or once you get a single aspect of it looking the way you want—copy the commands you used into a script. At the end of the process, you will have a file that can generate the same wonderful graphics at any time on any computer that runs Python.

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Start with the code at the beginning of Section 4.3.1 and write a script to produce a figure with a smooth, thick red line. Add labels for the axes, a title, and a legend. Make the text big enough to read easily.

4.3.4 Replacing curves

Sometimes you do not need a new figure—you may simply want to replot a modified version of your data. You can remove all data and formatting from the current figure by calling `plt.cla()`, PyPlot's clear axes command, and then calling `plt.plot` again. However, this removes all of your formatting.

To replace a curve without clearing all of your formatting, you can update the data of individual line objects within the plot:

```
plt.plot(x, 3*x)           # Plot two lines.
plt.plot(x, x**3)
plt.xlabel('Time [s]')      # Label axes.
plt.ylabel('Position [cm]')
5 ax = plt.gca()            # Assign a name to the current Axes object.
lines = ax.get_lines()      # Assign a name to its list of Line objects.
lines[1].set_data(x, x**2)  # Replace plot data of second line.
```

The shape of one curve will change, but all other properties of the plot will be preserved.

4.3.5 More about figures and their axes

PyPlot uses a hierarchy of objects to create a plot. The most important of these for our purposes are the `Figure` and `Axes` objects.

Calling `plt.figure()` creates and returns a `Figure` object. A common side effect of this function is to open a figure window on your screen. A `Figure` object is not capable of plotting a function; it simply contains and manages all of the elements of a plot. To actually make a graph, we need the second type of object: an `Axes` object.

An `Axes` object is always associated with a `Figure` object. We usually create an `Axes` object by calling PyPlot's convenient `plt.plot` function. This function adds an `Axes` object to the current figure

if none is present, uses it to graph the data we provide, and returns a list of the line objects that were added to the plot. The `Axes` object possesses all of the data and methods needed to draw a graph. Many of these methods create and manage yet more objects. You can access these subsidiary objects—things such as lines (associated with a `Line2D` object), axis labels (associated with Python strings), tick marks (associated with a NumPy `ndarray`), and so on—through the parent `Axes` object. Many of these manage their own subset of objects necessary to perform their function. You can assign a variable to any of these objects and control its behavior through the variable, as demonstrated in Section 4.3.4 above. This is often more convenient than accessing a property through the parent `Axes` object.

A full discussion of this topic lies well beyond the scope of this tutorial. Fortunately, PyPlot provides convenient tools for managing this hierarchy of objects without requiring much understanding of the details. Commands like `plt.plot`, `plt.xlim`, and `plt.legend` are convenience functions that use the methods of the current `Figure` and `Axes` objects to carry out common operations in making plots.

You can do a lot with PyPlot; however, to get the most control over your plots, you may need to delve into the details of Matplotlib's `Figure` and `Axes` objects. Exploring the objects returned by `plt.gcf()` (get current figure) and `plt.gca()` is a good place to start. The official documentation is at matplotlib.org.

4.3.6 T2 Error bars

To make a graph with error bars, use

```
plt.errorbar(x_values, y_values, yerr=y_errors, xerr=x_errors, fmt='or')
```

This function doesn't add error bars to an existing plot; rather, it creates a new plot (or a line within an existing plot) with error bars. Its syntax is similar to `plt.plot`, but it accepts additional optional arguments. Point `n` will be augmented by error bars that extend coordinate distance `y_errors[n]` above and below the point, and coordinate distance `x_errors[n]` to the left and right of the point. Both error arguments are optional. If you supply only one, there will be no error bars in the other direction. If you do not supply either, `plt.errorbar` behaves like `plt.plot`. To specify a marker for `errorbar`, you must use the keyword argument `fmt`. This example specifies red circles.

4.3.7 3D graphs

Sometimes a graph consists of points or a curve in a three-dimensional space. To make 3D plots, we must import an additional tool from one of the Matplotlib modules. We must also interact with PyPlot in a different way. The following script will plot a helix:

```
# line3d.py      [get code]
from mpl_toolkits.mplot3d import Axes3D      # Get 3D plotting tool.
fig = plt.figure()                          # Create a new figure.
ax = Axes3D(fig)                           # Create 3D plotter attached to figure.
5 t = np.linspace(0, 5*np.pi, 501)        # Define parameter for parametric plot.
ax.plot(np.cos(t), np.sin(t), t)           # Draw 3D plot.
```

Notice that this time we are using the `plot` method attached to a specific object. We created an `Axes3D` object and can now use its methods to generate three-dimensional plots. The method `ax.plot` accepts three arrays, whose corresponding entries are interpreted as the `x`, `y`, and `z` coordinates of each point

to be drawn. The object generated by `Axes3D(fig)` can also create surface plots, three-dimensional contour plots, three-dimensional histograms, and much more.

Your screen is two-dimensional. If you ask for a “3D” plot, what you actually get must be a two-dimensional projection—what a camera would see looking at your plot from some outside `viewpoint`. Switching to a different viewpoint may make it easier to see what’s going on. You may be able to click and drag within your plot to change the viewpoint. (This depends on the graphics back end you are using. See Section A.2.) If this is not possible, you can change the viewpoint from the IPython command prompt or in a script by issuing the command `ax.view_init(elev=30,azim=30)` prior to making the plot. (This command also works after making the plot, if you follow it by `plt.draw()`). You specify the viewpoint by giving the elevation and azimuth in degrees—not radians.

You can now label your plot axes by using `ax.set_zlabel('text')` and similarly for the others.

4.3.8 Multiple plots

Multiple plots on the same axes

You may have noticed in the previous examples that every new `plt.plot` command adds a new curve to an existing figure, if one is already open.

By default, a new plot is added to the current axes of the active figure.

You can also create several curves on the same axes with a single plot command. Try

```
x = np.linspace(0, 1, 51)
y1 = np.exp(x)
y2 = x**2
plt.plot(x, y1, x, y2)
```

This example shows that you can give `plot` more than one set of (x, y) pairs. Python will choose different colors for each curve, or you can specify them manually:

```
plt.plot(x, y1, 'r', x, y2, 'ko')
```

A third way to draw multiple curves on one set of axes is to give `plt.plot` an ordinary vector of x values, but a two-dimensional array of y values. Each *column* of y will be used to draw a separate curve using a common set of x values. This approach can be useful when you wish to look at a function for several values of a parameter.

Try the following code:

```
num_curves = 3
x = np.linspace(0, 1, 51)
y = np.zeros( (x.size, num_curves) )
for n in range(num_curves):
    y[:, n] = np.sin((n+1) * x * 2 * np.pi)
plt.plot(x, y)
```

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Add a legend explaining which curve is which, then use the methods of Section 4.3.3 to embellish the plot.

Multiple plot windows

If you want two or more separate plots open, use `plt.figure()` to create a new figure and make it active (the “current” figure), then use `plt.plot` to generate the second plot in the new figure.

If you don’t supply an argument, `plt.figure()` chooses a new figure number that is not in use yet, starting from 1 (not 0). You can also assign a name you choose yourself. For example, `plt.figure('Joe')` will create a figure named “Joe.”¹⁷ The next `plt.plot` command will draw its output in the current figure, with no effect on other figures.

Be aware that “Joe” is just a label for a figure, not a variable name assigned to the `Figure` object. You can create such a variable: `joe = plt.figure('Joe')`.

If you are generating several plots in a session, you will probably want to create a new figure each time by using `plt.figure()`. However, these plots can use a fair amount of your computing resources, and finding the graph you want among a dozen figure windows is not always simple. So once you are finished with a figure, you can close it. You can use `plt.close()` to close the current figure, or you can close a figure by using its label: `plt.close('Joe')`.

You can close all open figures with `plt.close('all')`.

You can include this command near the top of a script to ensure that all figures come from the latest run.

4.3.9 Subplots

You may wish to place multiple plots side by side in a single figure window for comparison. In Python jargon, you’d like multiple `axes` occupying the same `figure`. The function `plt.subplot(M, N, p)` divides the current figure window into a grid of cells with M rows and N columns, then makes cell number p the active one. Here, p must be an integer between 1 and M*N. (This is another case where Python does *not* start counting from 0.) Any graphing command issued now will affect cell number p. Try this example:

```
# subplot.py      [get code]
from numpy.random import random
t = np.linspace(0, 1, 101)
plt.figure()
5 plt.subplot(2, 2, 1); plt.hist(random(20))           # Upper left
plt.subplot(2, 2, 2); plt.plot(t, t**2, t, t**3 - t)   # Upper right
plt.subplot(2, 2, 3); plt.plot(random(20), random(20), 'r*') # Lower left
plt.subplot(2, 2, 4); plt.plot(t*np.cos(10*t), t*np.sin(10*t)) # Lower right
plt.suptitle("Data and Functions")                      # Overall
```

Each `plt.subplot` command selects a subregion of the figure. The subsequent `plt.plot` command sets up axes and draws a plot in the current subplot region. If your subplots do not fit nicely into the figure window or overlap one another, you can resize the figure window or try the command `plt.tight_layout()`.

Make sure that all your `plt.subplot` commands for a particular figure use the same values for N and M. If you change either, all existing subplots will be erased. For more freedom in placing subplots and

¹⁷ Or make it the current figure if it already exists. Either way, `plt.gcf()` returns the current `Figure` object, and `plt.gca()` returns the current `Axes` object of the current figure. So, you can use `plt.figure('Joe')` to select “Joe,” then `plt.gcf()` and `plt.gca()` to adjust this plot, even if there are multiple figures open.

insets, see `help(plt.axes)`. The `plt.suptitle` command at the end adds a centered title to the entire figure. If desired, `plt.title` can be used to add titles to the individual subplots.

PyPlot offers an alternate approach to creating and working with subplots—one with a similar name, but different behavior: `plt.subplots`. This command creates a new figure, lays out the subplots within, and returns both the `Figure` object and an array of `Axes` objects. Thus, it must be called when *creating* the figure, not after `plt.figure()`. The following code produces the same figure as the last example.

```
# subplots.py [get code]
from numpy.random import random
t = np.linspace(0, 1, 101)
fig, ax = plt.subplots(2,2)
ax[0,0].hist(random(20))          # Upper left
ax[0,1].plot(t, t**2, t, t**3 - t) # Upper right
ax[1,0].plot(random(20), random(20), 'r*') # Lower left
ax[1,1].plot(t*np.cos(10*t), t*np.sin(10*t)) # Lower right
fig.suptitle("Data and Functions")    # Overall
```

Note some of the differences compared to the previous example. First, we only use `plt` when creating the figure. We unpack and assign the objects it returns to the variables `fig` and `ax`, and then use their methods to do the rest of the plotting. Second, `ax` is an array of `Axes` objects. We access the individual subplots as we would access elements of any NumPy array.¹⁸ This means we do not have to use a special command to switch between subplots, as in the previous example.

You can use `plt.subplot` or `plt.subplots` to create the same figure. The official Matplotlib documentation encourages the “object-oriented” approach of `plt.subplots` over the “state-based” approach of `plt.subplot`, but the decision is largely a matter of personal preference. Just don’t mix the two, or you may encounter bizarre behavior.

4.3.10 Saving figures

All of your figures will disappear when you end your session in Spyder. If you used a script to create and adjust your plots, you can recreate them later, but you may also want to use plots in other applications or print them out. Saving a plot to a graphics file for such purposes is straightforward. The figure window has a `Save` icon that will open a dialog allowing you to save the current figure in a variety of formats. The default is `.png`, but most Python distributions will also allow you to save the figure in `.pdf`, `.jpg`, `.eps`, or `.tif` format. To get the complete list, consult the figure:

```
fig = plt.gcf()                      # Get current figure object.
fig.canvas.get_supported_filetypes()
```

All you need to do is give the figure a name with the appropriate extension; PyPlot will create a graphics file of the type you requested.

You can also save a figure from the IPython command prompt, or from within a script, by using the command `plt.savefig`. For example, to save the current figure as a `.pdf` file, type

```
plt.savefig("greatest_figure_ever.pdf")
```

¹⁸ If you call `plt.subplots` with no arguments or use it to create a figure with only one subplot, it will not return an array of `Axes` with a single element. It simply returns the `Figure` and a single `Axes` object.

The plot will be saved in your current working directory. (You can force Python to save in a different directory by giving the complete path, as discussed in Section 4.1.2, page 54.)

4.3.11 Using figures in other applications

If you save a figure as an `.eps`, `.pdf`, or `.svg` file, you can open and modify it in a *vector-graphics* application such as Inkscape or Xfig (both freeware), or a commercial alternative.¹⁹ Text such as the title or axis labels may be rendered as “outlines,” which can make it difficult to edit in another application. If you encounter this problem, you can instruct Python to save text as “type” in an `.svg` file. Before saving the figure, modify the parameter that controls how fonts are saved in `.svg` figures:

```
import matplotlib  
matplotlib.rcParams['svg.fonttype'] = 'none'
```

Then, save your figure as an `.svg` file.

Some applications won’t load these `.svg` files. An alternative that still gives “type” is to use

```
import matplotlib  
matplotlib.rcParams['pdf.fonttype'] = 42
```

Then, save your figure as a `.pdf` file, and edit it in another application. For more information about `matplotlib.rcParams`, see “Customizing matplotlib” at matplotlib.org/users/customizing.html.

If you are going to use figures in a presentation, you may instead need to save images in a *raster* format (also called **bitmap**), such as `.gif`, `.png`, `.jpg`, or `.tif`. For publication, however, vector graphics is generally best.

¹⁹ Inkscape: www.inkscape.org; Xfig: mcj.sourceforge.net/frm_installation.html.

CHAPTER 5

First Computer Lab

These exercises will use many ideas from the previous chapter. Our goals are to:

- Develop basic graphing skills;
- Import a data set; and
- Perform a simple fit of a model to data.

5.1 HIV EXAMPLE

Here we explore a model of the **viral load**—the number of virions in the blood of a patient infected with HIV—after the administration of an antiretroviral drug. One model for the viral load predicts that the concentration $V(t)$ of HIV in the blood at time t after the start of treatment will be

$$V(t) = A \exp(-\alpha t) + B \exp(-\beta t). \quad (5.1)$$

The four parameters A , α , B , and β are constants that control the behavior of the model. A and B specify the initial viral load, α is the rate at which new cells are infected, and β is the rate at which virions are removed from the blood.¹

In this lab, you will use what you have learned about Python up to this point to generate plots based on the model, import and plot experimental data, and then fit model parameters to the data.

5.1.1 Explore the model

To get started, launch Spyder, import NumPy and PyPlot, and then create an array by typing

```
time = np.linspace(0, 1, 11)
time
```

at the IPython command prompt. Press <Return> after each line. You should see a list of 11 numbers. Now modify these commands to create an array of 101 numbers ranging from 0 to 10 and assign it to the variable `time`.

Next, you will evaluate a compound expression involving this array by using the solution of the viral load model given in Equation 5.1.

The first step is to give the constants in the mathematical equation names you can type, such as `alpha` and `beta` instead of α and β . It is wise to give longer, more descriptive names even to the variables whose names you can type: for example, `time` for t and `viral_load` for $V(t)$. Now, set $B = 0$, and choose some interesting values for A , α , and β . Then, evaluate $V(t)$ by using the following command:

¹ Nelson, 2015, Chapter 1 gives details of the model and its solution.


```
viral_load = A * np.exp(-alpha*time) + B * np.exp(-beta*time)
```

You should now have two arrays of the same length, `time` and `viral_load`, so plot them:

```
plt.plot(time, viral_load)
```

Create a few more plots using different values of the four model parameters.

5.1.2 Fit experimental data

Now let's have a look at some experimental data.

Follow the instructions of Section 4.1 to obtain the data set `01HIVseries`. Copy files `README.txt`, `HIVseries.csv`, `HIVseries.npy`, and `HIVseries.npz` into your working directory, or be sure you can locate these files using paths. The `HIVseries` files contain time series data. Read the file `README.txt` for details.

Import the data set into Python by reading `HIVseries.csv` into an array or by loading either `HIVseries.npy` or `HIVseries.npz`. (Remember: The functions `np.load` and `np.loadtxt` return the data you requested, but you must assign that data to a variable to access it later.) You are free to give the data whatever name you like, but we will refer to the array as `hiv_data`.

If you use `np.loadtxt` to import `HIVseries.csv` or `np.load` to import `HIVseries.npy`, the data will be contained in a single array. Find the variable you created in the Variable Explorer. It has two columns of data. The first column is the time in days since treatment of an HIV-positive patient; the second is the concentration of virus in that patient's blood, in arbitrary units.

If you instead use `np.load` to import `HIVseries.npz`, the data will be contained in an object that does not show up in the Variable Explorer. (See Section 4.2.2.) You can nevertheless inspect its contents. Type `hiv_data.files` to find the names of the arrays stored within. There are two keys, called '`time_in_days`' and '`viral_load`'. You can inspect the corresponding arrays by asking for them by name. You can also assign the data in these arrays to a variable name of your choosing: for example, `concentration = hiv_data['viral_load']`.

Next, we are going to visualize the data. In order to plot the viral load as a function of time, you will need to separate the data into two arrays to pass to `plt.plot`. Do that and plot the data points now. Don't join the points by line segments. Instead, make each point a marker (symbol), for example, a small circle or plus sign. Label the axes of your plot. Give it a descriptive title, too. Such embellishments are discussed in Section 4.3.1.

Assignment:

- Plot the experimental data points and the function in Equation 5.1 on the same axes. Adjust the four parameters of the model until you can see both the data and the model in your plot.*

[Remark: You probably know about black-box software packages that do such “curve fitting” automatically. In this lab, you should do it manually, to see how the curves respond to changes in the parameters.]

The goal is now to tune the four parameters of Equation 5.1 until the model agrees with the data. It is hard to find the right needle in a four-dimensional haystack! We need a more systematic approach than just guessing. Consider the following: Assuming $\beta > \alpha$, how does the trial solution behave at long times? If the data also behave that way, can we use the long-time behavior to determine two of the four unknown constants, then hold them fixed while adjusting the other two?

Even two constants is a lot to adjust by hand, so let's think some more: How does the initial value $V(0)$ depend on the four constant parameters? Can you choose these constants in a way that always gives the correct long-time behavior and initial value?

- b. *Carry out this analysis so that you have only one remaining free parameter, which you can adjust fairly easily. Adjust this parameter until you like what you see.*
- c. *The latency period of HIV is about ten years, or 3,600 days. Based on your results, how does the inverse of the T-cell infection rate, $1/\alpha$, compare to the latency period? Is the latency period long because it takes HIV a long time to infect new cells? Or does the model suggest another scenario?*

5.2 BACTERIAL EXAMPLE

Now we turn our attention to genetic switching in bacteria. In 1957, A. Novick and M. Weiner studied the production of a protein called beta galactosidase in *E. coli* bacteria after introducing an *inducer* molecule called TMG.²

5.2.1 Explore the model

Here are two families of functions that come up in the analysis of the Novick–Weiner experiment:

$$V(t) = 1 - e^{-t/\tau} \quad \text{and} \quad W(t) = A \left(e^{-t/\tau} - 1 + \frac{t}{\tau} \right). \quad (5.2)$$

The parameters τ and A are constants.

Assignment:

- a. *Choose $A = 1$, $\tau = 1$, and plot $W(t)$ for $0 < t < 2$.*
- b. *Make several arrays $W1$, $W2$, $W3$, and so on, using different values of τ and A , and plot them all on the same axes.*
- c. *Change the colors and line styles (solid, dashed, and so on) of the lines.*
- d. *Add a legend to help a reader sort out the curves. Explore some of the other graph options.*

5.2.2 Fit experimental data

Follow the instructions of Section 4.1 to obtain the data set 15novick. Copy `g149novickA.csv`, `g149novickA.npy`, `g149novickB.csv`, `g149novickB.npy`, and `README.txt` into your working directory, or be sure you can locate these files using paths. The files contain time series data from a bacterial population in a culture.

Assignment:

[*Remark:* Again, you are asked not to use an automatic curve-fitting system. The method suggested in the Hints will give you a deeper understanding of the math than accepting whatever a black box spits out.]

² See Nelson, 2015, Chapter 10 for details of the experiment and model.

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- a. *Plot the experimental data points and your trial functions for $V(t)$ (see Equation 5.2) on the same axes, as you did for $W(t)$ before.*

When you plot the experimental data, do not join the points by line segments; make each point a symbol such as a small circle or plus sign. Label the axes of your plot. Select some reasonable values for the parameter τ in the model, and see if you can get a curve that fits the data well. Label the curves, then add a legend to identify which curve is which. [Hint: To find a good estimate of τ , make a semilog plot of the quantity $1.0 - \text{data}$ versus time, where `data` is the array of experimental data points. (Can you explain why this is helpful? See Section 4.3.2 for more information.)]

- b. *Now try the same thing using the data in `g149novickB.csv` or `g149novickB.npy`. This time throw away all the data with time value greater than ten hours, and attempt to fit the remaining data to the family of functions $W(t)$ in Equation 5.2.*

[Hints: (i) You can “throw away data” by slicing an array.

(ii) At large values of t (but smaller than ten hours), both the data and the function $W(t)$ become straight lines. Find the slope and the y intercept of the straight line determined by Equation 5.2 in terms of the two unknown quantities A and τ . Next, estimate the slope and y intercept of the straight line determined by the data. From this, figure out some good initial guesses for the values of A and τ . Then, tweak the values to get a nicer looking fit.]

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CHAPTER 6

Random Number Generation and Numerical Methods

*At its highest level, numerical analysis
is a mixture of science, art, and bar-room brawl.
— T. W. Koerner, *The Pleasures of Counting**

The preceding chapters have developed a basic set of techniques for importing, creating, and modeling data sets and visualizing the results. This chapter introduces additional techniques for exploring mathematical models and their predictions:

- Random numbers and Monte Carlo simulations
- Solutions of nonlinear equations of a single variable
- Solutions of linear systems of equations
- Numerical integration of functions
- Numerical solution of ordinary differential equations

In addition, this chapter introduces several new methods for visualizing data, including histograms, surface plots, contour plots, vector field plots, and streamlines.

We start with writing your own functions, an invaluable tool for exploring physical models.

6.1 WRITING YOUR OWN FUNCTIONS

Section 3.3.5 introduced a principle:

Don't duplicate. Define once, and reuse often.

In the context of parameters (fixed quantities), this means you should define a parameter's value just once at the start of your code and refer to it by name throughout the rest of the program. But code itself can contain duplications if we wish to do the same (or nearly the same) task many times. Just as with parameter values, you may later realize that something needs to be changed in your code. Changing every instance of a recurring code fragment can be tedious and prone to error. It is better to define a *function* once, then invoke it whenever needed. You may even use the same fragment of code in more than one of your scripts. If each script imports the same externally defined function, then changes that you make once in the function file will apply to all of your scripts.

Functions in Python can do almost anything. There are entire libraries of functions that can carry out mathematical operations, make plots, read and write files, and much more. Your own functions can do all of these things as well. Functions are ideal for writing code once and reusing it often.

6.1.1 Defining functions in Python

A function can be defined at the command prompt or in a file. The following example is a basic template for creating a function in Python:

```
# excerpt from measurements.py      [get code]
def taxicab(pointA, pointB):
    """
    Taxicab metric for computing distance between points A and B.
    5      pointA = (x1, y1)
           pointB = (x2, y2)
    Returns |x2-x1| + |y2-y1|. Distances are measured in city blocks.
    """
    10     interval = abs(pointB[0] - pointA[0]) + abs(pointB[1] - pointA[1])
           return interval
```

The “taxicab metric” determines the billable distance between points “as the cab drives” rather than “as the crow flies.” (That is, we use the total distance driven rather than the shortest straight line distance between the two points.)

The function consists of the following elements:

Declaration: The keyword `def` tells Python you are about to **define** a function. Function names must adhere to the same rules as variable names. (See Section 1.4.3.) It is wise to give your functions descriptive names. Remember: You will only have to define it once, but you will wish to reuse it often. Now is not the time to save typing by giving it a forgettable name like `f`.

Arguments: The name of the function is followed by the names of all its **arguments**: the data it requires to do its calculation.¹ In this case, `taxicab` requires two arguments. If it is called with only one argument or arguments of the wrong type, Python will raise a `TypeError` exception.

Colon: The colon after the list of arguments begins an indented block of code associated with the function. Notice the similarity with `for` and `while` loops and `if` statements. Everything from the colon to the end of the code block will be executed when the function is called.

Docstrings: The text between the pair of triple-quotes (" " " . . . " " ") is a **docstring**. It is a special comment Python will use if someone asks for `help` on your function, and it is the standard place to describe what the function does and what arguments are required. Python will not complain if you do not provide a docstring, but someone who uses your code might.

Body: The body of the function is the code that does something useful with the arguments. This example is a simple function, so its body consists of just two lines. More complicated functions can include loops, `if` statements, and calls to other functions, and may extend for many lines.

Return: In Python, a function always “returns” something to the program that invokes it. If you do not specify a return value, Python will supply an object called `None`. In this example, our function returns a `float` object.

¹ Arguments in the definition of a function are sometimes called “formal parameters.”

 Arguments in the definition of a function are sometimes called “formal parameters.”

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6.1 Writing your own functions 77

Now, let’s look at how a function call works. Enter the function definition for `taxicab` at the command prompt. (You may omit the docstring, but only this one time!) Then, type

```
fare_rate = 0.40          # Fare rate in dollars per city block
start = (1, 2)
stop = (4, 5)
trip_cost = taxicab(start, stop) * fare_rate
```

When the fourth line is executed, `taxicab` behaves like `np.sqrt` and other predefined functions (Section 1.4.2, page 14):

- First, Python assigns the variables in the function’s argument list to the objects passed as arguments.² It binds `pointA` to the same object as `start`, and `pointB` to the same object as `stop`. (See Appendix F for details.)
- Then, Python transfers control to `taxicab`.
- When `taxicab` is finished, Python substitutes the return value into the assignment statement for `trip_cost`.
- Python finishes evaluating the expression and assigns the answer to `trip_cost`.

Although we defined `start` and `stop` as tuples in this example, we can call our new function with any objects that understand what `thing[0]` and `thing[1]` mean: lists, tuples, or NumPy arrays.

Your
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Define a function to compute the straight-line distance between two points in three-dimensional space. Give it a descriptive name and an informative docstring. See what happens when you call it with the wrong number or type of arguments, and ensure that using `help` on your function will enable a user to diagnose and resolve the issue.

In some ways, a function is similar to a script. It is a fragment of code that is executed upon request. Unlike a script, a function is a Python object. It can be called by name (once it is defined or imported), and it can be called by another script or function. A function communicates with the calling program by way of its arguments and its return value. After evaluating the function, Python discards all of the function’s local variables, so remember:

If a function performs a calculation, `return` the result.

You can define functions at the IPython command prompt. However, if you plan to use a function more than once, you should save it in a file. You can place a single function in its own file such as `taxicab.py` and run this file as a script prior to using the function. (Running the file will define the function as if it had been entered at the command prompt.) You can also define a function within the script that uses it, as long as its definition gets executed before the function is first called.

If you will be using the same functions in multiple scripts and interactive sessions, you can create a **module** in which you define one or more functions in a single `.py` file. A module is a script that contains

²  Arguments passed to a function are sometimes called “actual parameters.”

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a collection of definitions and assignments. It can be imported into your session from the command prompt, or by placing an `import` command within a script. You can also import selectively, as described in Section 1.3.

Place `taxicab` and the function you wrote in Your Turn 6A (called, for example, `crow`) in a single file called `measurements.py` in your working directory. Then, type `import measurements`. You now have access to both functions under the names `measurements.taxicab` and `measurements.crow`. Type `help(measurements)` and `help(measurements.taxicab)` to see how Python uses the docstrings you provide. Note that if you give a *module* the same name as a *function* it contains, you still have to provide both the module and function name to Python. For example, if you save the `taxicab` function in a file called `taxicab.py` and then `import taxicab`, calling `taxicab(A,B)` will result in an error. To access the function, you must use `taxicab.taxicab(A,B)`.

Modules that you write and wish to use should be located in the same folder as your main script, which is probably the global working directory you specified during setup. (See Sections 4.1.1 and A.2.)³

6.1.2 Updating functions

If you have modified a function in a script or in a module that you created, you will need to instruct Python to use the newest versions. If the function is defined in a script, you can run the script again by using the [RUN▶] button (if the script is open in the Editor) or the `%run` command.

If your function is part of a module that you have imported, you may need to restart the IPython kernel or relaunch Spyder for the changes to take effect. Just typing `import my_module` again will *not* update the module—even after calling `%reset`. Alternatively, you can use a function called `reload` to update any module without restarting. This function is part of the `importlib` module.⁴ To access the `reload` function, type

```
from importlib import reload
```

For example, if you modify the `taxicab` function's code in `measurements.py` after importing `measurements` as a module, you need to save the file and then type the following command:

```
reload(measurements)
```

Again, calling `import` will only cause Python to load a module *if it has not already been imported*.

You must save a module and then use `reload` to update functions in a module you have already imported.

Reloading a module is usually necessary only in debugging. Once your functions are working properly, you need only import them once in a script or interactive session.

6.1.3 Arguments, keywords, and defaults

We have already seen that keyword arguments can modify the behavior of functions like `plt.plot`. You can also use keywords and give arguments default values in functions you write. The following example demonstrates both of these techniques:

³ If you know about “paths,” then you can create folders that Python can access from any working directory.

⁴ In Python 2.7, `reload` is a built-in function and does not need to be imported.


```
# excerpt from measurements.py      [get code]
def distance(pointA, pointB=(0, 0), metric='taxi'):
    """
    Return distance in city blocks between points A and B.
    If metric is 'taxi' (or omitted), use taxicab metric.
    Otherwise, use Euclidean distance.
    pointA = (x1, y1)
    pointB = (x2, y2)
    If pointB is omitted, use the origin.
    """
    if metric == 'taxi':
        interval = abs(pointB[0] - pointA[0]) + abs(pointB[1] - pointA[1])
    else:
        interval = np.sqrt( (pointB[0] - pointA[0])**2 \
                           + (pointB[1] - pointA[1])**2 )
    return interval
```

The `distance` function decides how to calculate the distance between points based on the value of `metric`. Both `pointB` and `metric` have default values. If values for these arguments are passed to the function, it will use them; if not, it will use the defaults. Run the script `measurements.py`, then try

```
distance( (3, 4) )
distance( (3, 4), (1, 2), 'euclid' )
distance( (3, 4), 'euclid' )                                # This is an error.
distance( pointB=(1, 2), metric='normal', pointA=(3, 4) )
```

The arguments to a function must either be in the correct order according to the function definition, as in the first two lines of this example, or be paired with a keyword, as in the last line. The third line results in an error because Python assigns the string literal '`'euclid'`', appearing here as a positional argument, to the variable `pointB`.

6.1.4 Return values

A function can take any type or number of arguments—or none at all. The return value of a function is a single object, but that object may be a number, an array, a string, or a collection of objects in a tuple or list. A function will return an object called `None` if no other return value is specified.

Suppose that you wish to write a function to rotate a two-dimensional vector. What should the arguments be? What should the function return?

The arguments should include the vector to be rotated and an angle of rotation. The function should return the rotated vector. Here is a function that will accomplish the task:

```
# rotate.py      [get code]
def rotate_vector(vector, angle):
    """
    Rotate a two-dimensional vector through given angle.
    vector = (x,y)
    angle = rotation angle in radians (counterclockwise)
    Returns the image of a vector under rotation as a NumPy array.
    """

```



```

10    rotation_matrix = np.array([[ np.cos(angle), -np.sin(angle) ],
                               [ np.sin(angle),  np.cos(angle) ]])
        return np.dot(rotation_matrix, vector)

```

This implementation of rotation creates a 2×2 matrix and then multiplies this matrix and the vector supplied as the first argument. The function does not modify the contents of `vector`. Instead, `np.dot` creates a new array, which is returned by the function.

Python allows you to **unpack** compound return values, that is, to assign the individual elements of an object to separate variables. Multiple assignments on a single line, such as `x, y = (1, 2)`, are a simple example of unpacking. Python can unpack any *iterable* object: a tuple, a list, a string, or an array.

There are several ways to unpack an object. The following calls to `rotate_vector` all work, but the components of the rotated vector are split up in different ways:

```

vec = [1, 1]
theta = np.pi/2
r = rotate_vector(vec, theta)
x, y = rotate_vector(vec, theta)
5 _, z = rotate_vector(vec, theta)
first, *rest = rotate_vector(vec, theta)

```

After executing these commands, you should find that `r` is a NumPy array with two elements. The other variables, `x`, `y`, and `z`, contain individual components of the rotated vector. The **underscore**, `_`, is a dummy variable whose value is discarded. (Underscore is a special variable, whose value is the result of the most recent command. We are using it to temporarily store a value that we do not need.) After the final line of this example, `first` contains the first element of the rotated vector and `rest` is a list that contains everything else.⁵ In this case, `rest` contains only one element, but if the function returned an array with 100 elements, `first` would still contain only the first value and `rest` would be a list (not a NumPy array) containing the other 99 elements. You could convert it to an array with `np.array(rest)`.

In addition to unpacking, you can also index or slice the return value of a function. For example, if you only need the second element of the rotated vector, you could type

```
w = rotate_vector(vec, theta)[1]
```

When evaluating this and similar expressions, Python first evaluates the function, then substitutes its return value in place of the function call.

6.1.5 Functional programming

Python offers programmers great flexibility in writing functions. We discuss some of these nuances in Appendix F. However, we strongly endorse the following guidelines for writing your own functions:

1. Pass data to a function only through its arguments.
2. Do not modify the arguments of a function.
3. Return the result of a calculation with a `return` statement.

Python allows you to circumvent all of these conventions, but your code will be easier to write, interpret, and debug if you adhere to them. Your functions will accept input (arguments) and produce output (a return value), without side effects.

⁵ This type of unpacking is not available in Python 2.

A **side effect** is any effect on the computer's state other than returning a value. Avoiding side effects is the cornerstone of *functional programming*. Sometimes side effects are useful, but consider the alternatives when writing your own functions.

Unintended side effects on arrays can be particularly troublesome. Python does *not* automatically create a local copy of an array inside a function. As a result, a function can modify the data in an array passed as an argument using array methods—including assignment of individual elements. (See Section 2.2.6, page 26.) Array methods like `x.sort()` and `x.fill(3)`, as well as in-place arithmetic like `x+=1` and `x*=2`, can also modify array data. Such side effects have the potential to speed up your code and reduce memory usage for operations on large arrays. You can **overwrite** the array and modify its elements without making any copies. However, this increase in performance comes at the cost of difficulty in debugging.

If you really do intend for a function to modify the elements of an array, you can still avoid side effects. Create a local copy of the array within the function, operate on the copy, and return the new array.⁶ Alternatively, you can create a placeholder array within the function, fill it with values, and return the new array. NumPy adheres to this principle: Functions like `np.cos(x)` and `np.exp(x)` return new arrays, with no effect on `x`. If you wish, you can replace the original array with the new array returned by the function in the main code, rather than inside the function.

The following function illustrates these principles of functional programming:

```
# average.py      [get code]
def running_average(x):
    """
    Return cumulative average of an array.
    """
    y = np.zeros(len(x))          # New array to store result
    current_sum = 0.0              # Running sum of elements of x
    for i in range(len(x)):
        current_sum += x[i]        # Increment sum.
        y[i] = current_sum / (i + 1.0) # Update running average.
    return y
```

The array being processed is passed as an argument. This array is not modified; the result of the calculation is returned in a new array.

If you need the extra performance and reduced memory usage that come from overwriting an array, use the information in Appendix F to plan your code carefully and avoid unintended consequences. However, try vectorizing your code using NumPy's efficient array operations before you resort to overwriting. Vectorized code will almost always run faster than loops you write yourself in Python.

6.2 RANDOM NUMBERS AND SIMULATION

There are many interesting problems in which we do not have complete knowledge of a system, but we do know the probabilities of the outcomes of simple events. For example, you know the probability of any given outcome in the roll of a single die is 1/6, but do you know how likely it is that the sum of a roll of

⁶ You can copy an array using the array's `copy()` method: `y = x.copy()`. After this statement, `x` and `y` will be independent arrays with the same data. Note that slicing does *not* create a copy of an array. See Appendix F.

five dice is less than 13? Rather than work out the combinatorics, you could roll five dice many times and determine the probability empirically.

A random number generator makes it possible for a computer to “roll dice” millions of times per second. Thus, you can use a random number generator to simulate a system described by a stochastic model in which the probability distributions of the parameters are known. You can determine the likely behavior of the system, even if you cannot work out the details analytically. Such calculations are often called “Monte Carlo simulations.”

6.2.1 Simulating coin flips

The simplest example of a stochastic system is a coin flip. Suppose that you want to simulate flipping a coin 100 times, record the number of heads or tails, and then repeat the whole series of 100 flips N times. This will generate a set of N numbers, each falling in the range of 0 to 100, inclusive.

How do we get Python to flip the coin for us? First, try typing `1>2` at the IPython console prompt, and then `2>1`. You’ll see that Python returns a Boolean value of `True` or `False` when it evaluates each of these expressions. You can simulate a coin flip by generating a uniformly distributed random number between 0 and 1, then checking to see if it is less than 0.5. If the comparison returns `True`, we record heads; if `False`, we record tails. Python can also use such values in numeric calculations: It converts `True` to 1 and `False` to 0.

The module `np.random` contains random number generators for several probability distributions. In this chapter, we will only need the “continuous uniform distribution” over the interval [0, 1). You should explore the other available probability distributions, too. To use these functions, we create a random number generator object, and then access its methods. To save typing, we will give the method we want the nickname `rand`:

```
rng = np.random.default_rng() # create a random number generator object
rand = rng.random             # assign its uniform distribution method to rand
```

To get a thousand random numbers, you can now say `rand(1000)`. In the future, if you wish to switch to a different probability distribution, you only need to change the second line of code above: Choose a different method of `rng` and assign it the same nickname `rand`. You can instead access random number generator functions directly from the `numpy.random` module, as we did in Section 1.3.2 and Section 4.3.9: `from numpy.random import random as rand`. This works, but the `default_rng` approach uses an algorithm for generating random numbers that is more efficient and has better statistical properties. Since we are going to be generating a *lot* of random numbers, we will use it from now on.

To make a series of independent flips, first create an array called `samples` that contains 100 random numbers generated by `rand`. (Consult `help(rand)` for a clue about how to do this easily.) You can then convert the random samples to a simulation of coin flips with `flips=(samples<0.5)`. Python evaluates the comparison item by item, and stores the result in `flips`. You can then count the number of heads by using `np.sum(flips)` or the array method `flips.sum()`. Repeat this several times to get a feel for how likely it is that exactly 50 heads occur in 100 flips of a fair coin.

6.2.2 Generating trajectories

We can adapt coin flipping to study random walks, Brownian motion, and a host of other interesting physical and biological systems. Let’s create a random walk of 500 steps. Our trajectory will consist of 500

x values and 500 *y* values. Following good practice (Section 3.3.5), begin with

```
num_steps = 500
```

The idea behind a random walk is that every step is a statistically independent, random event. You know from the preceding section how to get an array containing 500 random binary digits. Make two such arrays called *x_step* and *y_step*. For a coin flip, **True** and **False** or 1 and 0 were sufficient. For our random walk, however, we need a random string of +1 and -1 values.

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- a. Do a simple algebraic operation on *x_step* that maps each 1 to +1 and each 0 to -1. Do the same for *y_step*.
- b. Next, convert these arrays into successive positions of the random walker. Consult **help(np.cumsum)** to see why that function is useful.
- c. Write a script to draw your random walk. Run it several times.

Try writing a function called *get_trajectory* that will take the single argument *num_steps* and return the two arrays that contain the coordinates of a random walker.

6.3 HISTOGRAMS AND BAR GRAPHS

6.3.1 Creating histograms

A **histogram** is a graphical representation of a discrete probability distribution. Suppose that you wish to check that the random number generator *rand* really gives a uniform distribution. To make a simple histogram, try

```
# histogram.py [get code]
from numpy.random import random as rand
data = rand(100)
plt.hist(data)
```

A histogram appears, but a number of things have happened without your supervision. The function *plt.hist* has

- Inspected the array and determined its range;
- Divided that range into a number of equally spaced **bins**;
- Counted how many elements of *data* fall into each bin;
- Graphed the result as a set of bars; and
- Returned a tuple containing two arrays and a strange-looking item called <a list of 10 Patch objects>.

Consulting **help(plt.hist)** reveals the content of the return values as well as the keywords and defaults that you can use to control the output. For instance, there is an optional keyword argument called *bins* that specifies the number of bins into which data points will be sorted. The default behavior is to divide the entire range of data into 10 bins. However, if you have a few outliers, this can lead to a histogram

where most of your data falls into a single bin! You can use the keyword argument `bins='auto'` to ask

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Python to sample the data and determine an optimal bin size. This is a useful option if you are having trouble figuring out how many bins you should use.⁷

Another keyword argument controls the alignment of each bar: The default `align='mid'` specifies that the bars will be centered on the midpoint of each bin.

Compare the output when you specify 10, 100, and 1000 bins. For more control, you can provide the range over which you wish to bin the data using the `range` keyword, or you can specify the edges of each bin explicitly for nonuniform binning.

For control over presentation or for further analysis, you can get the counts and bin edges used to generate the plot by unpacking the return value of `plt.hist`.⁸

```
counts, bin_edges, _ = plt.hist(data)
```

The function returns three objects, so three variables are required to unpack its output. If you are interested only in the first two objects, you can assign the third to Python's dummy variable, named underscore, as shown here.

If you do not wish to generate a plot and only want the histogram data, you can use `np.histogram` instead. It is the same function that `plt.hist` uses to generate data for plotting:

```
counts, bin_edges = np.histogram(data)
```

Note that this function only returns two objects.

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Try it, and inspect the return values to make sure you understand how these functions generate a histogram. Note, in particular, the numbers of elements in the two arrays.

Once you've created the binned data, you can then plot it in another style or transform it prior to plotting. The `plt.hist` (or `np.histogram`) function has simply done the sorting and counting for you. For example, you can make a custom bar graph by using `plt.bar`. The arguments of this function are an array of bar positions, an array of bar heights, and a single width or an array of widths, which `plt.bar` uses to draw rectangles. All the arrays must be the same length, so you have to discard the last element of `bin_edges` returned by `plt.hist` or `np.histogram`, which is the right-hand edge of the last bin.

There are many ways to present data. The following code uses `plt.bar` to generate a colorful plot in which the width of a bar in the histogram is proportional to its height:

```
bin_size = bin_edges[1] - bin_edges[0]
new_widths = bin_size * counts / counts.max()
plt.bar(bin_edges[:-1], counts, width=new_widths, color=['r','g','b'])
```

⁷ An “optimal” bin size is one for which the histogram provides the best estimate of the underlying probability distribution. There are several methods that converge to the same result for very large data sets. See `help(np.histogram_bin_edges)` for a description of other methods available.

⁸ See Section 6.1.4 for unpacking, and Section 6.3.2 for more about bin edges.

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6.3.2 Finer control

As mentioned earlier, Python offers a lot of control over how your data is displayed. Sometimes you will wish to specify the edges of the bins or the range of values that each bin collects, not just the number of bins. Both `plt.hist` and `np.histogram` can accommodate this: Instead of passing an integer number of bins to either function, supply an array whose entries are the edges of the desired bins. You need to provide one more edge than the number of bins. (Each bin begins where the previous one ended, but you need to specify where the first one starts.) Be aware that Python will ignore any data points that fall outside the range you provide.

For example, to divide a collection of random numbers into bins that each span an inverse power of 2, such as $0, \frac{1}{128}, \frac{1}{64}, \frac{1}{32}, \dots, \frac{1}{2}, 1$, you can use `np.logspace` to generate the bins:

```
log2bins = np.logspace(-8, 0, num=9, base=2)
log2bins[0] = 0.0 # Set first bin edge to zero instead of 1/256.
plt.hist(data, bins=log2bins)
```

If you bin the data by using `np.histogram` and then do some additional analysis, you can use `plt.bar` to display it as a histogram. The following code will do this:

```
bin_size = bin_edges[1] - bin_edges[0]
plt.bar(bin_edges[:-1], counts, width=bin_size, align='edge')
```

By default, the `plt.bar` function draws each bar centered on the corresponding value in the first argument. This behavior is generally not appropriate for a histogram. The preceding code fragment specifies a different behavior with the `align` keyword argument. (Try omitting it to see the difference.)

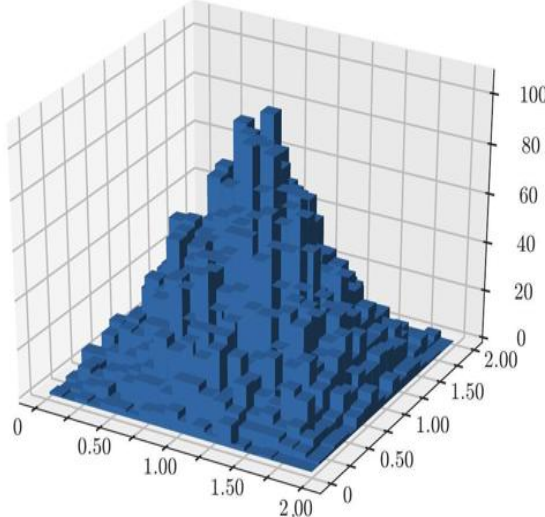


Figure 6.1: A 3D bar plot (“Lego plot”) made by `Axes3D.bar3d`.

It is also possible to create histograms in higher dimensions, in which data are binned along two or more different axes. NumPy provides `np.histogram2d` to bin a collection of xy pairs. You can display the

result as a three-dimensional bar chart by using `Axes3D.bar3d` (Figure 6.1), or as a grid of colored pixels (a heat map) by using `plt.imshow`. (Section 8.1.1, page 109 gives an example.) `np.histogramdd` extends binning to any number of dimensions.

6.4 CONTOUR PLOTS, SURFACE PLOTS, AND HEAT MAPS

In earlier chapters, we saw several methods for plotting data sets with a single independent variable, such as a time series. Such data lend themselves to two-dimensional plots. However, models with two or more independent variables require higher dimensional plots.

A function of two variables, $h(x, y)$, can be interpreted as a surface whose height over each point (x, y) has been specified, just as Earth's topography is specified by the altitude at every latitude and longitude. Useful graphical representations of the height are contour plots, surface plots, and heat maps. A contour plot is a two-dimensional drawing in which contour lines are used to represent the height, as in a topographic map; a surface plot is a 3D perspective drawing of the surface itself; a heat map instead uses color to indicate height.

6.4.1 Generating a grid of points

In order to make a plot of $h(x, y)$, you must specify the heights at a finite set of points. Typically you'll want those points to form a grid in the xy plane. Python gives a convenient way to construct such a grid: First, set up a 1D array of x values covering the desired range and an analogous 1D array of y values. Then, call the function `np.meshgrid` to create the grid. If the first array has N entries and the second has M entries, then `np.meshgrid` will return two $M \times N$ arrays. The arrays contain the x and y coordinates of each grid point, respectively. Try

```
x_vals = np.linspace(-3, 3, 21)
y_vals = np.linspace(0, 10, 11)
X, Y = np.meshgrid(x_vals, y_vals)
```

Inspect `X` and `Y` to make sure you understand the result: `np.meshgrid` returned a list containing two arrays, which we unpacked and assigned to `X` and `Y`. Each is itself an array with $11 \times 21 = 231$ entries giving the x and y coordinates of the grid points.⁹ You can now evaluate a function on these arrays, perhaps by using vectorized math, to produce a third array called `Z`, and then create contour plots, surface plots, or heat maps using `(X, Y, Z)`. Sometimes it is useful to create the intermediate array of distances, `R=np.sqrt(X**2+Y**2)`, if the height only depends on the distance from the origin.

6.4.2 Contour plots

Suppose that we wish to visualize the function $z(x, y) = \cos x \sin y$. Creating a contour plot is simple using the arrays `X` and `Y` generated above:

```
Z = np.cos(X) * np.sin(Y)
plt.contour(X, Y, Z)
```

⁹ Python sets `X[i, j]=x_vals[j]` and `Y[i, j]=y_vals[i]`. You can change this convention with the keyword argument `indexing='ij'`, which will set `X[i, j]=x_vals[i]` and `Y[i, j]=y_vals[j]` (the transposes of the default behavior).

The number of contour lines is 10 by default. To change this, add a fourth argument to the function call: `plt.contour(X, Y, Z, 20)`. Instead of an integer, you can also pass an array that contains the exact “heights” (z values) for which you want to see contours. You can control the appearance of contour lines by using keywords, and you can even label the contour lines with the following commands:

```
# contour.py      [get code]
cs = plt.contour(X, Y, Z, 10, linewidths=3, colors='k')
plt.clabel(cs, fontsize=10)
```

The `plt.contour` function returns a `ContourSet` object, and the `plt.clabel` command knows how to add labels to the contours in this object.

The `plt.contour` function draws only contour lines. A related function called `plt.contourf` will draw filled contours. You can change the set of colors used by PyPlot using either the `cmap` keyword argument or the command `plt.set_cmap`. If you enter the name of a nonexistent color map, Python will return an error that lists all of the available color maps.

6.4.3 Surface plots

Creating a surface plot is similar to creating a contour plot. The arguments to the functions are the same, but PyPlot has to access functions from a different module in order to create three-dimensional graphics. Recall Section 4.3.7:

```
from mpl_toolkits.mplot3d import Axes3D      # Import 3D plotting tool.
```

An `Axes3D` object can draw a variety of three-dimensional plots. For example, the following commands will create a surface plot:

```
ax = Axes3D( plt.figure() )      # Create 3D plotter attached to new figure.
ax.plot_surface(X, Y, Z)        # Generate 3D plot.
```

It takes a lot of computational resources to create a three-dimensional plot, so Python tries to use a small number of points when generating a surface. Its default is to use up to 50 grid points in each direction. If your arrays contain more points, Python will omit some of them. To use all of your points, supply the keyword arguments `rstride` and `cstride` (“row stride” and “column stride”) with the value 1:

```
ax.plot_surface(X, Y, Z, rstride=1, cstride=1)
```

You can also increase resolution by specifying the total (or maximum) number of points to use along each direction. Use the keyword arguments `rcount` and `ccount` (“row count” and “column count”):

```
ax.plot_surface(X, Y, Z, rcount=100, ccount=200)
```

will create a 100 by 200 point grid for drawing the surface (20 000 points total).

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Make a surface plot of the function $z = x^2 + y^2$ over a suitable grid of values, where x and y range from -1 to 1 . Use keyword arguments to create a low-resolution surface and to use every data point.

The default of `plot_surface` is to use a single color and shading from imaginary sources of light to render a surface. You can get a more colorful plot by supplying a color map with the `cmap` keyword.¹⁰ (`plt.set_cmap` has no effect on surface plots.)

6.4.4 Heat maps

Creating a heat map is similar to creating a contour plot or a surface plot. The arguments to the functions are the same, but PyPlot uses color to indicate height, rather than contour lines or a three-dimensional projection. PyPlot's `pcolormesh` function is a fast and useful tool for creating heat maps of a surface.¹¹ Try the following command:

```
plt.pcolormesh(X, Y, Z)
```

You should see a grainy color image of the function. The default behavior of `plt.pcolormesh` is to plot each cell of the image with a single color. However, a look at `help(plt.pcolormesh)` reveals a way to get a smoother image by interpolating the color between points in our array: use the optional `shading` keyword argument. Compare the output of the previous command with that of

```
plt.pcolormesh(X, Y, Z, shading='gouraud')
```

As with a contour plot, you can change the color scheme by supplying a color map with the `cmap` keyword or by using `plt.set_cmap` after plotting.

On color figures

Graphs, contour plots, surface plots, and heat maps look great in color. However, colors on the screen don't always translate well to printouts. If you will be printing an assignment in grayscale or publishing a paper in a print journal, you may get better results with a colorless color map like '`gray`' or '`bone`'.

Also keep in mind that color can convey information in a figure, but only if the viewer knows what the colors mean. To help the viewer, you can easily add a colorbar—a legend for the color scheme—to a plot with `plt.colorbar`, or use the `colorbar` method of a figure. All three types of plot—contour, surface, and heat map—require an extra step to add a colorbar: You must assign a variable to the object returned by the plotting command, and then pass this object to `colorbar`. The following commands illustrate how to add a colorbar:

```
plt.figure()
contours = plt.contour(X, Y, Z, cmap='jet')
plt.colorbar(contours)

5 fig = plt.figure()
ax = Axes3D(fig)
surface = ax.plot_surface(X, Y, Z, cmap='coolwarm')
fig.colorbar(surface)

10 plt.figure()
heatmap = plt.pcolormesh(X, Y, Z, cmap='bone')
plt.colorbar(heatmap)
```

¹⁰ In the blog that accompanies this book, we describe other options for coloring and shading 3D surfaces. (See press.princeton.edu/titles/32489.html.)

¹¹ Section 8.2 will discuss another method that uses image processing tools.

Finally, keep in mind that up to 8% of your male readers may be unable to discriminate red from green, the most common form of “colorblindness.” When possible, add words or redundant graphical information to improve accessibility, instead of relying entirely on color.

6.5 NUMERICAL SOLUTION OF NONLINEAR EQUATIONS

It is often necessary to solve nonlinear equations when studying physical and biological systems. For example, determining the fixed points in a single-gene toggle system requires finding the *roots* of a sixth-order polynomial.¹² There are no general analytic solutions for polynomials of degree greater than four, and even for cubic equations the formulas are cumbersome. Biological oscillators may involve still more complicated functions that are not even polynomials. Numerical methods for finding the roots of such expressions are quite useful.

6.5.1 General real functions

The methods developed to find the roots of functions are closely related to methods for optimizing functions. Neither Python nor NumPy has a collection of optimization tools, but SciPy has an extensive library called `scipy.optimize`.¹³ You can import the entire module:

```
from scipy import optimize
```

The command `dir(optimize)` will tell you the names of all of the functions available in this module, and `help(optimize)` will provide the details. However, we will not need most of the functions in this library, so we will simply import individual functions as they are required.

A useful function for finding roots is called `fsolve`. It takes two arguments: a function name and a point (or array of points) at which `fsolve` begins searching for roots.

You can define the function to be solved by using the approach described in Section 6.1. If you need to solve a relation like $g(x) = 7$, define your function as $f(x) = g(x) - 7$. The `fsolve` function determines the zeroes of whatever function you provide.

While `fsolve` is quite good at finding roots, it will do even better if you help it. You can provide an array of points *near* the zeros of a function, then have `fsolve` refine these. A good way to generate initial guesses is to plot the function and estimate visually where it crosses the x axis.

Example: Try the following and explain the results:

```
from scipy.optimize import fsolve
def f(x): return x**2 - 1
fsolve(f, 0.5)
fsolve(f, -0.5)
fsolve(f, [-0.5, 0.5])
```

Solution: The equation $x^2 - 1 = 0$ has two roots, but which root `fsolve` returns depends on where we tell the function to begin searching. Providing an array of starting points yields an array of roots found from those starting points.

¹² A root is a zero of a function: a value of x for which $f(x) = 0$.

¹³  Its functions are adapted from the highly optimized MINPACK library of FORTRAN functions.

Example: Now try the following, and explain what you see:

```
def f(x): return np.sin(x)**10
fsolve(np.sin, 1)
fsolve(f, 1)
```

Solution: The last expression above does not return exactly 0. What you are seeing is an example of *numerical error*, caused by the finite number of digits that Python uses to represent a number. The result is a number that is close the correct solution, but not exact.

The phrase “numerical error” is standard, but unfortunate. You didn’t make any error; neither did Python. Your computer’s hardware simply has limited precision.

A more serious type of error can occur when searching for the roots of a function with a *singularity*.¹⁴ Inspect the output of `fsolve(f, 2)` when $f(x) = 1/(x - 1)$. This function has a singularity at $x = 1$ and never crosses the x axis. However, `fsolve` may return a solution without raising any errors or warnings. We can learn more by using a keyword to instruct `fsolve` to provide more information:

```
def f(x): return 1/(x-1)
fsolve(f, 2, full_output=True)
```

This time, the value of the root is returned along with an object that contains a lot more information. You can refer to `help(fsolve)` to learn what the terms mean, but the last line is a sure sign of trouble: The function reached its maximum number of evaluations without satisfying its requirements for a root. If your numerical result doesn’t make sense, set `full_output=True` and consult `help(fsolve)`.

6.5.2 Complex roots of polynomials

The solutions returned by `fsolve` are restricted to be real numbers. Consider the equation $1/x = 1 + x^3$. In order to use `fsolve`, we must manipulate this into $f(x) = x(1 + x^3) - 1$ and solve for the roots. Assume that we have some good reason to guess that the real roots are near 1 and -1 :

```
def f(x): return x * (1 + x**3) - 1
fsolve(f, 1)
fsolve(f, -1)
```

A quartic equation has four roots; but, in the present case, `fsolve` can only find two of them because the other two are complex.

You can find all of the roots—real or complex—of any *polynomial* by using the NumPy function `np.roots`. This function takes a vector of coefficients that define a polynomial (see `help(np.roots)` for details). You can find all of the roots of the polynomial above with `np.roots([1, 0, 0, 1, -1])`.

T2 Complex nonlinear equations

For arbitrary nonlinear equations (those that are not polynomials) with complex roots, a straightforward application of `fsolve` or `np.roots` may not be sufficient. For example, the function $f(z) = \frac{1}{z} - (1 + z^2)^4$ has one real and two complex roots. `fsolve` can easily find the real root, but if you need the other two, you will have to work harder.

¹⁴ A singularity is a point where a function is not mathematically well defined or well behaved. Common examples are $1/x$ (division by zero), $|x|$ (not differentiable), or $\sin(x)/x$ (indeterminate form) evaluated at $x = 0$.

For a function of one variable, `fsolve` searches along the real axis. However, it is capable of finding numerical solutions to N equations in N unknowns. For a function of a complex variable, we can find complex zeros by solving two equations in two unknowns. The two equations are the real and imaginary parts of the function, and the two unknowns are the real and imaginary parts of the complex variable. The following code gives all three roots of $f(z)$:

```
def f(x):
    z = x[0] + 1j*x[1]
    q = 1/z - 1 - z**2.4
    return q.real, q.imag
5
fsolve(f, [2, 0])  # Finds real root.
fsolve(f, [0, 2])  # Finds first complex root.
fsolve(f, [0,-2])  # Finds second complex root.
```

We can also solve this particular equation by using `np.roots`. First, we can rewrite $f(x) = 0$ as $x^{3.4} = 1 - x$ and raise both sides to the fifth power to generate a 17th-order polynomial. Then, we can get all 17 roots using `np.roots`. However, most of these solutions are extraneous, so we can use a `for` loop to substitute them into the original expression and discard the spurious solutions.

```
# Create monomial with NumPy's polynomial class.
X = np.poly1d([1,0])

# Create polynomial whose roots might be solutions to the original problem.
5 P = X**17 - (1-X)**5

# Examine roots and ignore extraneous solutions.
for z in np.roots(P):
    if np.isclose(1/z - z**2.4 - 1, 0): print(z)
```

Using `np.roots`—or computer algebra packages like Wolfram Alpha or SymPy (see Section 10.3)—gives exact results for simple problems, but such programs are easily overwhelmed. For instance, solving the almost identical function $g(z) = \frac{1}{z} - (z^{2.41} + 1)$ exceeds the allowed computation time on Wolfram Alpha, and numerical error in `np.roots` makes the zeros of the 341st degree polynomial useless for the original problem. The strictly numerical approach using `fsolve` encounters no such difficulty.

6.6 SOLVING SYSTEMS OF LINEAR EQUATIONS

Often we need to solve a set of simultaneous linear equations. For example, a least-squares fit of a two-parameter linear model to a data set involves solving equations of the general form

$$\begin{bmatrix} a_1 \\ a_2 \end{bmatrix} = \begin{bmatrix} c_{11} & c_{12} \\ c_{21} & c_{22} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix}.$$

We are given (or can find) the a 's and c 's, and wish to know the x 's.

Python and NumPy do not provide a collection of linear algebra tools, but SciPy provides a library for matrix mathematics called `scipy.linalg`.¹⁵ To gain access to the functions in this module, we need to

¹⁵ Its functions are adapted from the highly optimized LAPACK library of FORTRAN functions.

import them. You can import the entire module.

```
from scipy import linalg
```

To learn more about the module and the functions it provides, use `dir(linalg)` and `help(linalg)`.

Some of the more commonly used functions are

<code>inv</code>	matrix inverse
<code>det</code>	determinant
<code>sqrtm</code>	matrix square root
<code>expm</code>	matrix exponentiation
<code>eig</code>	eigenvalues and eigenvectors of a matrix
<code>eigh</code>	eigenvalues and eigenvectors of a Hermitian matrix
<code>svd</code>	singular value decomposition

We will import these functions by name as they are needed, rather than importing the entire library.

To solve the linear system above, we can compute the inverse of the matrix C and multiply the vector a by this matrix. The order of multiplication is important: The solution of $a = C \cdot x$ is obtained by multiplying both sides *on the left* by C^{-1} .

```
# matrix_inversion.py [get code]
from scipy.linalg import inv
a = np.array([-1, 5])
C = np.array([[1, 3], [3, 4]])
x = np.dot(inv(C), a)
```

This puts the solution to the problem into the array x , as you can check by computing `np.dot(C, x)-a`. (Again, the result may not be exactly zero, due to numerical error. However, the difference should be on the order of 10^{-15} or less.)

6.7 NUMERICAL INTEGRATION

Integrating a function is a common task in physical modeling. For example, you may wish to integrate a probability density function to determine the probability that its variable will fall within a certain range.

As with linear algebra, Python and NumPy do not support numerical integration, but SciPy does. The appropriate module is called `scipy.integrate`.

```
from scipy import integrate
```

The command `dir(integrate)` will tell you the names of all of the functions in this module, and `help(integrate)` will provide more information. However, we will not need most of the functions in the library, so we will simply import individual functions as they are required.

There are several integration routines available in the module, each with particular strengths. The all-purpose workhorse is called `quad`, from “quadrature,” an old word for integration.¹⁶

6.7.1 Integrating a predefined function

To carry out a numerical integration, you must provide a function name for the integrand and limits of integration. Optional arguments provide more control of the integration process.

¹⁶ This function is adapted from the highly optimized QUADPACK library of FORTRAN functions.

To see how this works, let's evaluate

$$\int_0^{x_{\max}} dx \cos(x)$$

for various values of x_{\max} . Try this code:

```
# quadrature.py      [get code]
from scipy.integrate import quad
x_max = np.linspace(0, 3*np.pi, 16)
integral = np.zeros(x_max.size)
5 for i in range(x_max.size):
    integral[i], error = quad(np.cos, 0, x_max[i])
plt.plot(x_max, integral)
```

Here is how the code operates:

- The second line imports the `quad` function.
- The third line creates an array of values for `x_max`.
- The fourth line sets up an array of the same shape as `x_max` to store the results.
- The next two lines carry out the integrations and store the results. By default, `quad` returns two values: the result of the integral and an estimate of the error. Line 6 unpacks these values. The value of the error is overwritten during each iteration, but each result gets saved in its own slot of an array.

The `quad` function evaluates `np.cos` at a carefully selected series of points and uses the result to approximate the integral over the range given by its second and third arguments.

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Do the integral by hand, and check whether `quad` got it right.

6.7.2 Integrating your own function

Next, suppose that you wish to integrate a function that is not predefined. If your function is short, you can define it at the command prompt:

```
def f(x): return x**2
```

If the function is long and complicated, or if you plan on using it more than once, you'll want to define it in a script or module, as described in Section 6.1. Once you have defined your function, you can pass its name (with no arguments or parentheses) to `quad` and integrate it like any other function.

Be aware that `quad` will only integrate real functions. To integrate a complex function, integrate its real and imaginary parts separately. If you give `quad` a function that returns complex values, it issues a warning but does not halt your program.

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- Integrate $f(x) = x^2$ from 0 to 2, and check that `quad` gets it right.
- Try a function whose integral you *don't* know: Evaluate

$$\int_0^{x_{\max}} dx e^{-x^2/2}$$

for values of x_{\max} from 0 to 5, and plot the results.

- Can `quad` handle infinite limits? Use `-np.inf` and `np.inf` as limits to evaluate

$$\int_{-\infty}^{+\infty} dx e^{-x^2/2}.$$

Compare to the exact result: $\sqrt{2\pi}$.

6.7.3 Oscillatory integrands

Sometimes we wish to evaluate an integral whose integrand oscillates rapidly. This can cause `quad` to fail to converge using its default settings. However, `quad` is an adaptive method, and if you allow it to subdivide the integration range sufficiently, it will generally yield satisfactory results. The keyword `limit` controls how fine a grid `quad` is allowed to use. Try

```
quad(np.cos, 0, 5000)           # Results in a warning and enormous error!
quad(np.cos, 0, 5000, limit=1000) # No warning; accurate result
np.sin(5000)                   # Exact result for comparison
```

6.7.4 Parameter dependence

The `quad` function can only integrate functions of a single variable. If you have a function of several variables and you wish to integrate one of them, holding the others fixed, there are two options: You can define a dummy function of one variable and pass the dummy function to `quad`, or you can supply a keyword and value to `quad` that will specify the constant arguments. For example, suppose that you have defined a function:

```
def f(x, a, b, c): return a*x**2 + b*x + c
```

To integrate the function over x holding $(a, b, c) = (1, 2, 3)$, the two options work as follows:


```
# Use dummy function.
def g(x): return f(x, 1, 2, 3)
integral1, err = quad(g, -1, 1)
# Use keyword.
5 integral2, err = quad(f, -1, 1, args=(1, 2, 3))
```

If the variable you wish to integrate is not the *first* argument of the function, you cannot use the second method. You must define a dummy function.

6.8 NUMERICAL SOLUTION OF DIFFERENTIAL EQUATIONS

In the physical and life sciences, it is often possible to write down a system of differential equations that govern a system or describe the behavior of a model. However, it may be impossible to solve this system in terms of known functions. A classic example is the three-body problem in classical mechanics: $\mathbf{F} = m\mathbf{a}$ and Newton's law of gravitation are all that is required to write down the differential equations, but no one in the last four centuries has been able to find a general solution!

Numerical solution is a powerful tool in studying such systems.¹⁷ Starting from some initial configuration of a system, you can calculate the next configuration from the given differential equations. From *that* configuration, you can calculate the *next* configuration by again using the equations, and so on. Computers are ideal for executing this repetitive operation, and efficient libraries have been developed for the task.

To solve differential equations, we turn once again to SciPy. The module we need is called `scipy.integrate`, the same module that contains `quad`. There are several functions available in the module, but we will restrict our attention to the one called `odeint`.¹⁸

```
from scipy.integrate import odeint
```

6.8.1 Reformulating the problem

An ordinary differential equation (ODE) is one that describes a function of a single independent variable, which we'll call $x(t)$. A textbook example is the driven harmonic oscillator:

$$\frac{d^2x}{dt^2} = -x + g(t).$$

The `odeint` function can "only" solve ODEs of the form

$$\frac{dy}{dt} = \mathbf{F}(\mathbf{y}, t), \quad (6.1)$$

where \mathbf{y} is a vector whose components y_i are functions of t , and \mathbf{F} is a vector whose components are functions of y_i and t . Fortunately, *any* explicit ODE can be put into this form. For example, to describe the

¹⁷ Many authors use the phrase "numerical integration of an ODE" to denote the procedure we call "numerical solution of an ODE." We will reserve the word "integration" for actually performing integrals, a procedure that sometimes, but not always, can be used to solve ODEs.

¹⁸ This function is adapted from the highly optimized ODEPACK library of FORTRAN functions. Calling `odeint` invokes a routine called LSODA, an adaptive solver that chooses a predictor-corrector method or a backward differentiation formula depending on its progress with the problem.

second-order ODE for the driven harmonic oscillator to `odeint`, we must reformulate it as a system of coupled first-order equations. First, define two new variables that will be the components of the vector \mathbf{y} :

$$y_1 = x \quad y_2 = \frac{dx}{dt}.$$

Next, write the derivatives of y_1 and y_2 in terms of y_1 , y_2 , and t :

$$\frac{dy_1}{dt} = \frac{dx}{dt} = y_2 \quad \frac{dy_2}{dt} = \frac{d^2x}{dt^2} = -x + g = -y_1 + g.$$

This allows us to cast a second-order differential equation in the form required by `odeint` (Equation 6.1): \mathbf{y} is an array with two entries, and $\mathbf{F}(\mathbf{y}, t)$ is a function that returns an array with two entries:

$$\mathbf{y} = \begin{bmatrix} y_1 \\ y_2 \end{bmatrix} \quad \frac{d\mathbf{y}}{dt} = \mathbf{F}(\mathbf{y}, t) = \begin{bmatrix} y_2 \\ -y_1 + g \end{bmatrix}. \quad (6.2)$$

Most common ODEs can be recast in a similar manner.

6.8.2 Solving an ODE

You must supply three arguments to `odeint`: the name of the function $\mathbf{F}(\mathbf{y}, t)$, an array $\mathbf{y}(t_0)$ that defines the initial conditions, and an array of t values at which you'd like `odeint` to evaluate the solution $\mathbf{y}(t)$. The general function call is

```
y = odeint(F, y0, t)
```

The variables in the expression are:

F – A function $\mathbf{F}(\mathbf{y}, t)$ that accepts a 1D array and a scalar and returns an array.

y0 – A one-dimensional array with the initial values of \mathbf{y} .

t – An array of t values at which \mathbf{y} is to be computed. The first entry of this array is the time at which the initial values $y0$ apply.

y – An array of the values of $\mathbf{y}(t)$ at the points specified in **t**.

To determine the motion of an undriven harmonic oscillator (the example above with $g(t) = 0$), we first need to define the function $\mathbf{F}(\mathbf{y}, t)$. The first argument, \mathbf{y} , is an array that contains the values of y_1 and y_2 at time t . The function we need is given in Equation 6.2 with $g = 0$:

```
# simple_oscillator.py      [get code]
def F(y, t):
    """
    Return derivatives for second-order ODE y'' = -y.
    """
    dy = [0, 0]          # Create a list to store derivatives.
    dy[0] = y[1]          # Store first derivative of y(t).
    dy[1] = -y[0]         # Store second derivative of y(t).
    return dy
```


Note that `odeint` requires $F(y, t)$ to accept time as an argument, even though the value of t is not used in this example. You will get an error if you omit this argument in the definition of $F(y, t)$.

We can now study the motion of the harmonic oscillator for different initial conditions:

```
# solve_ode.py      [get code]
"""ODE solver for harmonic oscillator."""

import numpy as np, matplotlib.pyplot as plt
from scipy.integrate import odeint

# Import ODE to integrate:
from simple_oscillator import F

10 # Create array of time values to study:
t_min = 0; t_max = 10; dt = 0.1
t = np.arange(t_min, t_max+dt, dt)

# Provide two sets of initial conditions:
15 initial_conditions = [ (1.0, 0.0), (0.0, 1.0) ]

plt.figure()    # Create figure; add plots later.
for y0 in initial_conditions:
    y = odeint(F, y0, t)
    plt.plot(t, y[:, 0], linewidth=2)

    skip = 5
    t_test = t[::skip]                      # Compare at a subset of points.
    plt.plot(t_test, np.cos(t_test), 'bo')    # Exact solution for y0 = (1,0)
25 plt.plot(t_test, np.sin(t_test), 'ro')    # Exact solution for y0 = (0,1)
```

This example illustrates the following:

- Like `quad`, `odeint` expects its first argument to be a function name (with no parentheses).
- The second argument is a vector with two entries, specifying the initial conditions.
- The return value of `odeint` is an array. The first column is $y(t)$ evaluated at the values in t ; the second is dy/dt . If you solve a higher order ODE, the third column will contain d^2y/dt^2 , and so on.

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Modify `simple_oscillator.py` to incorporate the driving force $g(t) = \sin(0.8t)$, and comment on how the solution changes.

6.8.3 T2 Parameter dependence

If we wanted to explore several different harmonic oscillators, we might consider modifying the function $F(y, t)$ as follows:


```

# parametric_oscillator.py      [get code]
def F(y, t, spring_constant=1.0, mass=1.0):
    """
    Return derivatives for harmonic oscillator:
    y'' = -(k/m) * y
    y = displacement in [m]
    k = spring_constant in [N/m]
    m = mass in [kg]
    """
    dy = [0, 0]          # Array to store derivatives
    dy[0] = y[1]
    dy[1] = -(spring_constant / mass) * y[0]
    return dy

```

To specify fixed parameters when using `odeint`, there are two options, just as for `quad` in Section 6.7.4. First, we can define a dummy function that sets the parameters and pass this function to `odeint`. Alternatively, we can supply an optional argument to `odeint` by using the keyword `args`. Suppose that the system has a spring constant of 2.0 N/m and a mass of 0.5 kg.

```

y0 = (1.0, 0.0)
t = np.linspace(0, 10, 101)
k = 2.0
m = 0.5
# Use dummy function.
def G(y, t): return F(y, t, k, m)
yA = odeint(G, y0, t)

# Use keywords.
yB = odeint(F, y0, t, args=(k, m))

```

To use the second method, the parameters you wish to set must come after `y` and `t` in the list of arguments to `F`. Otherwise, you must define a dummy function.

6.8.4 Other ODE solvers

`odeint` is a fast, simple, and reliable function. However, there are many methods for numerical solution of ODEs, and `odeint` only provides access to one of them. Another function in the `scipy.integrate` library provides a single interface to several numerical methods: `solve_ivp`.

The input and output of `solve_ivp` differ substantially from `odeint`. The following example illustrates the use of `solve_ivp` for the simple harmonic oscillator.


```

# ivp_comparison.py      [get code]
""" Compare different ODE solvers using solve_ivp. """
import numpy as np, matplotlib.pyplot as plt
from scipy.integrate import solve_ivp
5
# Define ODE to integrate: simple harmonic oscillator.
def f(t,y): return [ y[1], -y[0] ]

# Define time interval.
10 t_min = 0
t_max = 10

# Define initial conditions.
y0 = [1.0, 0.0]
15

# Integrate the ODE using defaults.
result = solve_ivp(f, (t_min, t_max), y0)
plt.plot(result.t, result.y[0], '^k', label='RK45')

20 # Specify time series and use different solver.
dt = 0.1
t_vals = np.arange(t_min, t_max + dt, dt)

result = solve_ivp(f, (t_min, t_max), y0, t_eval=t_vals, method='BDF')
25 plt.plot(result.t, result.y[0], '.r', label='BDF')

plt.legend()

```

Note the following differences between `solve_ivp` and `odeint`.

- The order of function arguments is reversed. `solve_ivp` requires functions of the form `f(t,y)`, while `odeint` requires functions of the form `F(y,t)`. To use the functions defined in this chapter with `solve_ivp`, you can define a dummy function:

```

from simple_oscillator import F
def f(t,y): return F(y,t)

```

- `solve_ivp` only requires the initial and final times. It will automatically generate an array of intermediate times. You can specify the intermediate times with the optional `t_eval` keyword argument.
- An optional `method` keyword allows you to select different numerical methods. The default is '`RK45`', a fourth-order Runge-Kutta method, similar to MATLAB's `ode45` routine. Calling `solve_ivp` with `method='LSODA'` uses the same method as `odeint`. See `help(solve_ivp)` for more options.
- The result is packaged in a complex object, not a single array. This example accessed the time and position arrays through `results.t` and `results.y[0]`, respectively.

If your objective is to find a solution to a set of ordinary differential equations, `odeint` is a simple and reliable first option. If `odeint` gives perplexing results or if you wish to compare different numerical methods, try `solve_ivp`.

6.9 VECTOR FIELDS AND STREAMLINES

A vector field is a function whose value at any point in space is a vector. Common examples from physics include the electric field, the gravitational field, and the velocity field of a fluid. PyPlot provides two useful functions for visualizing vector fields and their streamlines: `plt.quiver` and `plt.streamplot`.

6.9.1 Vector fields

You can plot a two-dimensional vector field with `plt.quiver`. This function requires four arguments, all of which are arrays of the same size. The first two arguments define a grid of (x, y) values. (See Section 6.4.3.) Instead of specifying a scalar function like height or temperature at these points, the next two arguments specify the components of a two-dimensional vector \mathbf{v} at the corresponding (x, y) coordinates. Python will draw an arrow starting at each grid point to represent the vector field $\mathbf{v}(x, y)$.

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Try the following:

```
# vortex.py      [get code]
coords = np.linspace(-1, 1, 11)
X, Y = np.meshgrid(coords, coords)
Vx, Vy = Y, -X
5 plt.quiver(X, Y, Vx, Vy, pivot='mid', angles='xy')
plt.axis('square')
```

Explain the “vortex” pattern you get. (The keyword argument `mid` places center of each arrow at its grid point. The keyword argument `angles='xy'` ensures that the arrows will accurately represent the gradient of a function—like a velocity field in a fluid or an electric field.)

Python’s default arrows may not be what you need. `help(plt.quiver)` reveals several options for controlling the appearance of the arrows. For example, if you want your arrows to be the length that you specified in your arrays, the keyword arguments `units='xy'` and `scale=1` will produce this behavior. If the components of the vector are $(1, 0)$ at some point, the corresponding arrow will have a length of 1.0 in the scale used to draw the axes. (This means the arrows will get larger as you zoom in and smaller as you zoom out.)

Vector fields often arise as the gradient of a scalar function. For example, gravitational fields and electric fields are gradients of scalar potentials, and Fick’s law states that the flux of particles at a point is proportional to the gradient of the concentration:

$$\mathbf{J} = -D \nabla c.$$

NumPy can estimate the gradient of a function evaluated on a grid via `np.gradient`.¹⁹ The following code displays the gradient of a two-dimensional bell curve on its contour plot. Consult `help(np.gradient)` to explore its arguments and return value.

¹⁹ An analytic formula for the gradient of a function is usually more accurate than `np.gradient`, but such formulas are not available for experimental measurements.

```

# gradient.py      [get code]
import numpy as np, matplotlib.pyplot as plt

coords = np.linspace(-2, 2, 101)
5 skip = 5
X, Y = np.meshgrid(coords[::skip], coords[::skip]) # Coarse grid for gradient
R = np.sqrt(X**2 + Y**2)
Z = np.exp(-R**2)
x, y = np.meshgrid(coords, coords)                  # Fine grid for contour plot
10 r = np.sqrt(x**2 + y**2)
z = np.exp(-r**2)

ds = coords[skip] - coords[0]                         # Coarse grid spacing
dX, dY = np.gradient(z, ds)                          # Calculate gradient.

15 plt.contourf(x, y, z, 25)
plt.set_cmap('coolwarm')
plt.quiver(X, Y, dX.T, dY.T, scale=25, angles='xy', color='k')
plt.axis('equal')

```

Because of the way NumPy calculates the gradient and our convention for x and y axes, we must take the *transpose* of the arrays that `np.gradient` returns—that is, we must interchange rows and columns.

Python is also capable of drawing three-dimensional quiver plots using the `quiver3d` method of an `Axes3D` object. You must supply three arrays of coordinates and three arrays of vector components: `ax.quiver3d(X, Y, Z, Vx, Vy, Vz)`. You can use this function to visualize complex three-dimensional flow patterns or electromagnetic fields.

6.9.2 Streamlines

A system of first-order ordinary differential equations defines a vector field. One way to find solutions is to follow the arrows, generating “streamlines.” The word comes from an analogy to water flow: The velocity of the water defines a vector field; the trajectory of any small volume of water is a streamline. Electric and magnetic “field lines” are also streamlines of electric and magnetic fields.

Python’s `plt.streamplot` function generates several trajectories from a vector field. The syntax is

```
plt.streamplot(x, y, Vx, Vy)
```

You can adjust the length and density of lines by using optional arguments. See `help(plt.streamplot)` for details. The four arguments are the same as those used in `plt.quiver`. They specify the coordinate grid and the vector field to follow.²⁰

To illustrate the use of this function, begin with the vector field you drew in Section 6.9.1:

```

# streamlines.py      [get code]
lower, upper, step = -2, 2, 0.1
coords = np.arange(lower, upper + step, step)
X, Y = np.meshgrid(coords, coords)

```

²⁰ The documentation for `plt.streamplot` specifies 1D arrays for x and y ; however, passing 2D arrays generated by `meshgrid`

works just as well.

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```
5 vx, vy = y, -x
plt.streamplot(X, Y, vx, vy, linewidth=2)
plt.axis('square')
```

Notice that we chose a finer grid of points than before. Such a fine grid would have made a cluttered picture of the vector field with too many arrows. For generating streamlines, a fine grid yields more accurate results. (Python doesn't need to interpolate as much as it would with a coarse grid.)

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- Perhaps the picture drawn in the preceding example was a bit too predictable. Replace line 5 with

```
vx = y - 0.1 * x
vy = -x - 0.1 * y
```

and explain the streamlines that you find.

- Next, replace line 5 with

```
vx, vy = x, -y
```

and explain what happens.

You can fine-tune a streamline plot by using keyword arguments: `density` controls the number of lines in the plot. `start_points` allows you to specify the points the streamlines will pass through with a list of (x, y) pairs. `integration_direction` specifies which way to integrate from those starting points. `minlength` allows you to eliminate short streamlines. With some experimentation, you can generally find parameters to produce a very nice plot.

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CHAPTER 7

Second Computer Lab

In this lab, you'll use Python to generate two-dimensional random walks, plot their trajectories, and look at the distribution of end points for a large number of random walkers. You will also use numerical experiments to study the statistics of rare events. Our goals are to:

- Generate random walk trajectories that begin at the origin and take random diagonal steps:

$$x_{n+1} = x_n \pm 1 \quad y_{n+1} = y_n \pm 1. \quad (7.1)$$

- Plot individual trajectories side by side and compute statistics for many trajectories.
- Generate a Poisson distribution and analyze a Poisson process.

7.1 GENERATING AND PLOTTING TRAJECTORIES

First, review Section 6.2.

Our first task is to create a random walk of 1000 steps, each given by Equation 7.1.¹ Each trajectory will be a list of 1000 x values and 1000 y values. It's good programming practice to define the size of the simulation at the beginning of a script:

```
num_steps = 1000
```

Now you can set the size of all arrays by using `num_steps`.

Assignment:

- a. Use the ideas in Section 6.2 to make a random walk trajectory, and then plot it. To remove any distortion, use the command `plt.axis('square')` or `plt.axis('equal')` after making the plot.
- b. Now make four such trajectories, and look at all four side by side. Use `plt.figure()` to create a new figure window. You can access the individual subplots by using commands like `plt.subplot(2,2,1)` before the first `plt.plot` command, `plt.subplot(2,2,2)` before the second `plt.plot` command, and so on. Python may give each plot a different magnification. Consult `help(plt.xlim)` and `help(plt.axis)` to find out how to give each of your plots the same x and y limits.²

¹ Section 6.2.2 (page 82) introduced random walks.

² You can also use `plt.subplots` with the keyword arguments `sharex` and `sharey` (Section 4.3.9, page 68).

7.2 PLOTTING THE DISPLACEMENT DISTRIBUTION

Run your script several times, and compare the resulting trajectories. Your plots always look different! Sometimes the walker wanders off the screen; sometimes it remains near the origin. And yet, there is some family resemblance among them. Let us begin to understand in what sense they are similar.

How far does the random walker get after 1000 steps? More precisely, what is the distance from the starting point $(0, 0)$ to the ending point (x_{1000}, y_{1000}) for each of many random walks? We can use Python to give a statistical answer to such questions. Instead of four walks, we now need many, say, 100. You could manually examine all 100 plots, but it would be hard to see the common features. Instead, we'll ask Python to generate all of these random walks, but show us only a summary.

One straightforward way to make 100 walks is to take the code you already wrote and embed it inside of a `for` loop. You could create three arrays, `x_final`, `y_final`, and `displacement`, to store the ending `x` and `y` positions and distance to the origin. Then, just before the end of the loop, you could add something like

```
x_final[i] = x[-1]
y_final[i] = y[-1]
displacement[i] = np.sqrt(x[-1]**2 + y[-1]**2)
```

(You can also solve the problem by using vectorized operations instead of a `for` loop. Try to figure out how. The vectorized approach is faster than a loop if the arrays are not too large—that is, if `num_walks*num_steps` is smaller than 10^7 .)

We can summarize the results in at least three ways. You have a lot of end points (`x_final`, `y_final` pairs), so you can make a *scatter plot* by using `plt.plot` or `plt.scatter`. Alternatively, you can examine the lengths of the final displacement vectors, or their squares.

Assignment:

- a. Once you have a code that works, increase the number of random walks from 100 to 1000. (See Section 3.3.5.) Make a scatter plot of the end points.
- b. Use `plt.hist` to make a histogram of the displacement values.
- c. Make a histogram of the quantity `displacement**2`.
- d. Your result from (c) may inspire a guess as to the mathematical form of the histogram. Use semilog and log-log axes to test for exponential or power law relationships.
- e. Use `np.mean` to find the average value of `displacement**2` (the mean-square displacement) for a random walk of 1000 steps.
- f. Find the mean-square displacement of a 4000-step walk. If you wish to carry the analysis further, see if you can determine how the mean-square displacement depends on the number of steps in a random walk.

It turns out that random walks are partially predictable after all. Out of all the randomness comes regular *statistical* behavior, partly visible in your answers to (b–f).

Experimental data also agree with these predictions. The random walk, although stripped of much of the complexity of real Brownian motion, nevertheless captures nontrivial aspects of Nature that are not

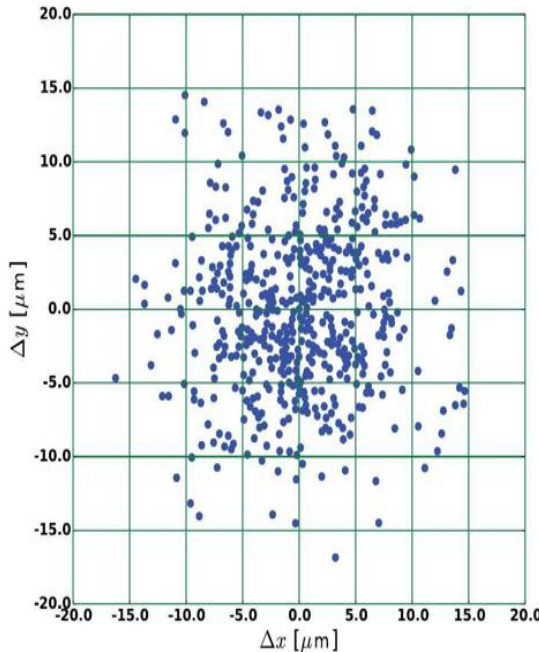


Figure 7.1: Experimental data for Brownian motion. Each dot is the final position after a fixed amount of time for a particle that started at the center of the figure.

self-evident from its formulation. See if your output qualitatively resembles the experimental data shown in Figure 7.1 for the diffusion of a micrometer-size particle in water.

7.3 RARE EVENTS

7.3.1 The Poisson distribution

Imagine an extremely unfair coin that lands heads with probability ξ equal to 0.08 (not 0.5). Each trial consists of flipping the coin 100 times. You might expect that we'd then get “about 8” heads in each trial, although we could, in principle, get as few as 0, or as many as 100.

The **Poisson distribution** is a discrete probability distribution that applies to rare events.³ For our extremely unfair coin, the Poisson distribution predicts that the probability of the coin coming up heads ℓ times in 100 flips is

$$\mathcal{P}(\ell) = \frac{e^{-8} \cdot 8^\ell}{\ell!}, \quad (7.2)$$

where ℓ is an integer greater than or equal to 0.

³ Nelson, 2015, Chapter 4 discusses this distribution.

Assignment:

- a. Before you start flipping coins, plot this function for some interesting range of ℓ values. You may find the following helpful:
 - The `factorial` function can be imported from `scipy.special`.
 - You need not take ℓ all the way out to infinity. You'll see that $P(\ell)$ quickly gets negligibly small.
 - In Python, the elements of a vector are always numbered 0, 1, 2, 3, ..., and ℓ is also an integer starting from zero, so ℓ is a good array index.
 - The value of 8^ℓ can get very large—larger than the largest integer NumPy can store.⁴

To avoid numerical *overflow*—and erroneous results—use an array of floats instead of integers. Consult `help(np.arange)`, and read about the `dtype` keyword argument.
- b. Perform N coin flip trials, each consisting of 100 flips of a coin that lands heads only 8% of the time. [Good practice: Eventually you may want to take N to be a huge number. While developing your code, make it not so huge, say, $N = 1000$, so that your code will run fast.]
- c. Get Python to count the number of heads, M , for each trial. Then, use `plt.hist` to create a histogram of the frequency of getting M heads in N trials. If you don't like what you see, consult `help(plt.hist)`. (For example, `plt.hist` may make a poor choice about how to bin the data.)
- d. Make a graph of the Poisson distribution (Equation 7.2 above) multiplied by N . What's the most probable outcome? Graph this plot on the same axes as the histogram in (c).
- e. Repeat (b-d) for $N = 1\,000\,000$, and comment. (This may take a while.)

Run the script over and over for $N = 1000$, and observe that the distribution is a bit different every time, and yet each plot has a general similarity to the others.

7.3.2 Waiting times

If we flip our imagined coin once every second, then our string of heads and tails becomes a time series called a **Poisson process**, or *shot noise*. Flipping heads is a rare event, because $\xi = 0.08$. We expect long strings of tails, punctuated by occasional heads. This raises an interesting question: After we get a heads, how many flips go by before we get the next heads? More precisely, what's the *distribution* of the *waiting times* from one heads to the next?

Here's one way to use Python to answer that question. We can make a long list of ones and zeros, then search it for each occurrence of a 1 with NumPy's `np.nonzero` function. This function takes an array of numbers and returns an array of the indices of its nonzero elements. Consult `help(np.nonzero)` and experiment with small arrays like `np.nonzero([1, 0, 0, -1])` to understand its behavior.

Each waiting time is the length of a run of zeros, plus one. You can subtract successive entries in the array returned by `np.nonzero` to find the waiting time between successive heads, then make a graph showing the frequencies of those intervals. NumPy's `np.diff` function will take the difference of successive entries in an array. Flatten the array returned by `np.diff` before plotting. See Section 2.2.8 for details on flattening arrays. Consult `help(np.diff)` for more information about that function.

Try to guess what this distribution will look like before you compute the answer. Someone might reason as follows: "Because heads is a rare outcome, once we get a tails we're likely to get a lot of them

⁴ By default, NumPy uses 64-bit integers, so the largest number it can store is $2^{63} - 1$.

in a row, so short strings of zeros will be less probable than medium-long strings. But eventually we're bound to get a heads, so *very* long strings of zeros are also less common than medium-long strings." Think about it. Is this sound reasoning? Now, compute the distribution. If your output is not what you expected, try to figure out why.

Assignment:

- a. *Construct a random string of 1's and 0's representing 1000 flips of the unfair coin. Then, plot the frequencies of waiting times of length 0, 1, 2, ..., as outlined above. Also make semilog and log-log plots of these frequencies. Is this distribution a familiar-looking function?*
- b. *What is the average waiting time between heads?*
- c. *Repeat (a) and (b) for 1 000 000 flips of the coin.*

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CHAPTER 8

Images and Animation

*The commonality between science and art is in trying to see profoundly
—to develop strategies of seeing and showing.*
— Edward Tufte

This short chapter introduces a few more useful tools for physical modeling with Python. First, we introduce tools for image processing. Then, we describe how to create animations and videos.

8.1 IMAGE PROCESSING

A digital image is a collection of pixels. The image can be stored in a variety of formats. In the case of a black-and-white photograph, each pixel is represented by a number corresponding to the intensity (mean photon rate) sensed at a point in the *xy* plane of the detector.¹ A digital camera takes the light intensity in each pixel and reports it as an integer between 0 and $2^m - 1$, where m is called the *bit depth*. A common choice is $m = 8$ (256 distinct light levels). Thus, a grayscale image can be represented by a single array of 8-bit integers, often called a “channel.” A color image from a digital camera usually consists of *three* such arrays—one each for **red**, **green**, and **blue** light in the *RGB color scheme*. Other digital images may include a fourth channel, called **alpha**, that controls transparency (*RGBA color scheme*).

A computer uses the pixel data in an image file to reproduce the image on your monitor. As far as Python is concerned, then, *every image is an array of numbers*. Conversely, *any array of numbers can be displayed as an image*. This mapping between arrays and images allows us to use Python to import, analyze, transform, and save images.

8.1.1 Images as NumPy arrays

Let’s use Python to import and display an image file. We first need to give Python access to some image data. Follow the instructions of Section 4.1 (page 53) to obtain the data set `16catphoto`. Copy the file `bwCat.tif` into your working directory.

Neither basic Python nor NumPy contains modules for working with images, but PyPlot contains functions for reading, displaying, and saving image files: `plt.imread`, `plt.imshow`, and `plt.imsave`. Import the photograph to an array with the following command:

```
photo = plt.imread('bwCat.tif')
```

¹ We are discussing raster images, also called bitmaps, which are saved in formats like `.png`, `.tif`, or `.jpg`. Another category, called vector graphics, represents figures mathematically in formats like `.svg` or `.eps`; see Section 4.3.10.

(If you have trouble loading image files, you may need to install the `pillow` library. See Section A.1.2 for details.) Verify that the photograph is indeed represented as an array of numbers by looking at some elements of `photo`. You should see the array in the Variable Explorer. You can also inspect the data fields of the array or examine a slice:

```
photo.shape
photo.dtype
photo[:10, :10]
```

This photograph is represented as a 648×864 array of integers. The `data type` of the array is `uint8`, which means that each value is an `unsigned integer` represented by 8 bits, matching the bit depth of the original image.

8.1.2 Saving and displaying images

We can now use PyPlot to display the array as an image:

```
plt.imshow(photo)
```

The resulting figure is probably not what you expected. PyPlot's defaults are convenient for plotting mathematical functions, but not black and white photographs. The following commands will set the color map, remove the axes, and change the background color for more convenient viewing.

```
plt.set_cmap('gray')          # Use grayscale for black and white image.
plt.axis('off')              # Get rid of axes and tick marks.
fig = plt.gcf()              # Get current figure object.
fig.set_facecolor('white')    # Set background color to white.
```

Now we see a recognizable image of a cat. You may find it convenient to define a single function to perform all these steps and display a black and white image from an array.

You can export any figure in a photographic format by using the `SAVE` button in the figure window. You can also create an image file from the command prompt or within a script:

```
plt.imsave('cat.jpg', photo, cmap='gray')
```

The function `plt.imsave` treats the array in the same way as `plt.imshow`, and it will use the current color map. Thus, we need to specify an appropriate color map for a black and white image.

8.1.3 Manipulating images

Because Python converts an image into a numerical array, you can perform mathematical operations on the array, potentially enhancing the image.

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Try this:

```
new_cat = (photo < photo.mean())
```

Display the new array, and explain what you see. Compare `new_cat` and `photo` in the Variable Explorer.

In Chapter 9, you'll learn about some common techniques for manipulating and enhancing images. However, you should be aware of the limits imposed by scientific ethics on modifying images to be used as evidence. (See Cromey, 2010.)

8.2 DISPLAYING DATA AS AN IMAGE

It is often useful to display a two-dimensional data set as an image. One way is to create a heat map using `plt.pcolormesh`, as described in Section 6.4.4. However, if you wish to represent each data point by a single pixel in an image, or if you have several arrays that could be interpreted as channels in an RGB image, then you can use the image processing tools of PyPlot.

Sometimes an image can reveal structure in a matrix or a data set. `plt.imshow` will create an image from a two-dimensional array. Try the following code:

```
M = np.zeros((40,40))
M[:10,:10] = np.random.random((10,10))
M[10:,:10:] = np.random.random((30,30))
plt.imshow(M)
5 plt.colorbar(_)
```

We can see the “block diagonal” structure of the matrix without inspecting a table of numbers.

Notice that with `plt.imshow`, points on the screen follow the conventional location of points in a matrix or a spreadsheet:

(0,0) is in the upper left corner (first column and first row); row number increases from top to bottom; and column number increases from left to right.

This is fine when displaying matrices or photos, but it is not the usual convention for graphing functions:

Cartesian coordinates place the origin at the lower left; x -values increase as you move to the right; and y -values increase as you move up.

If your data consists of samples of a function of position, $f(x,y)$, you'll probably want to place the origin in the lower left corner. You can use the keyword argument `origin="lower"` to accomplish this.

The preceding paragraph also implies another conflict between mathematical conventions and image conventions: For an array, the row index comes first and the column index comes second. But for Cartesian coordinates, x -values correspond to column indices and y -values to row indices. Hence, you may need to transpose your array when plotting it with `plt.imshow`. (If you use `np.meshgrid` and vector math to create your function, then you do *not* need to take the transpose.)

The following example illustrates these points:

```
# data_images.py      [get code]
""" Illustrate differences in image and Cartesian coordinates. """
import numpy as np, matplotlib.pyplot as plt

5 # Define a coordinate grid.
# Same spacing, but twice as many points along x-axis
x_max, y_max = 2, 1
```

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```

10 # Create coordinate arrays.
x = np.linspace(0,x_max,x_num)
y = np.linspace(0,y_max,y_num)

# Assign function values to placeholder array.
15 z = np.zeros((x_num,y_num))
for i in range(x_num):
    for j in range(y_num):
        z[i][j] = (x[i] - 2*y[j])**2

20 # Use meshgrid to generate the same function values.
X,Y = np.meshgrid(x,y)
Z = (X-2*Y)**2

# Visualize results.
25 fig, ax = plt.subplots(2,3, figsize=(12,6))
fig.suptitle(r"Plots of $f(x,y) = (x-2y)^2$")

ax[0,0].imshow(z)
ax[0,0].set_title("Loop: Image Coordinates")
30
ax[0,1].imshow(z, origin="lower")
ax[0,1].set_title("Loop: Spatial Coordinates")

ax[0,2].imshow(Z)
ax[0,2].set_title("meshgrid: Image Coordinates")

35 ax[1,0].imshow(z.transpose(), origin="lower")
ax[1,0].set_title("Loop: Transpose + Spatial Coordinates")

40 ax[1,1].imshow(Z, origin="lower")
ax[1,1].set_title("meshgrid: Spatial Coordinates")

ax[1,2].pcolormesh(X, Y, Z)
ax[1,2].axis('image')
45 ax[1,2].set_title("pcolormesh")

```

The three images along the top row do not follow the usual mathematical conventions; the three images along the bottom row do. Notice that `plt.pcolormesh` uses coordinate values to label the axes—not array indices.

In summary,

Array indices are not Cartesian coordinates.

If you wish to visualize a matrix or data that has no spatial meaning, `plt.imshow` is a simple choice. If you are plotting a function $f(x,y)$, it is usually simpler to use `np.meshgrid` and `plt.pcolormesh`.

to create a heat map.

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8.3 ANIMATION

A picture may be worth a thousand words, but a motion picture can be even better. Matplotlib contains a module called `animation` for creating movies from plots. We can also view a collection of still plots (*frames*) in sequence to create a simple animation.

We provide two scripts to create animations below. They illustrate two different approaches, but they have a common design principle: Create an empty plot, take control of the line or point objects you wish to animate, then update them in each frame. Once you understand how to do this, you can create a variety of animations.

8.3.1 Creating animations

The following script will create a movie of one of the random walks you studied in Chapter 7 by using the function `FuncAnimation` from the `animation` module. Use Python's `help` function to explore the many options available.

```
# walker.py      [get code]
# Jesse M. Kinder -- 2021
""" Make a movie out of the steps of a two-dimensional random walk. """
import numpy as np, matplotlib.pyplot as plt
5 from matplotlib.animation import FuncAnimation

# Create a random number generator.
rng = np.random.default_rng() # create a random number generator object
rand = rng.random             # assign its uniform distribution method to rand
10
# Set number of steps for each random walk.
num_steps = 100

# Create an empty figure of the desired size.
15 plt.close('all')           # Clear anything left over from prior runs.
bound = 20
fig = plt.figure()            # Must have figure object for movie.
ax = plt.axes(xlim=(-bound, bound), ylim=(-bound, bound))

20 # Create empty line and point objects with no data.
# They will be updated during each frame of the animation.
my_line, = ax.plot([], [], lw=2)          # Line to show path
my_point, = ax.plot([], [], 'ro', ms=9)    # Dot to show current position

25 # Generate the random walk data.
x_steps = 2*(rand(num_steps) < 0.5) - 1  # Generate random steps: +/- 1.
y_steps = 2*(rand(num_steps) < 0.5) - 1
x_coordinate = x_steps.cumsum()           # Sum steps to get position.
y_coordinate = y_steps.cumsum()

30 # This function will generate each frame of the animation.
# It adds all of the data through frame n to a line
```

```

# and moves a point to the nth position of the walk.
def get_step(n, x, y, this_line, this_point):
    35   this_line.set_data(x[:n+1], y[:n+1])
    this_point.set_data(x[n], y[n])
    return this_line, this_point

# Call the animator and create the movie.
40 my_movie = FuncAnimation(fig, get_step, frames=num_steps,
                           fargs=(x_coordinate, y_coordinate, my_line, my_point) )

# Save the movie in the current directory.
# *** NEXT LINE WILL CAUSE AN ERROR UNLESS FFMPEG IS INSTALLED. ***
45 # my_movie.save('random_walk.mp4', fps=30, dpi=300)

```

In lines 22 and 23, `ax.plot` returns a list with a single object. We need the object inside the list, not the list itself. The commas after `my_line` and `my_point` force Python to unpack this list rather than assign the variable to the list.

This script uses a new approach to plotting to improve video quality: Fixed elements of the plot—axes, tick marks, legends, labels, and so on—are not changed from one frame to the next. Instead of drawing each frame “from scratch,” the script creates a figure and axes just once (lines 17–18). Then, it creates two variables and assigns them to line and point objects that initially contain no data (lines 22–23).² The function `get_step` modifies the data of the line and point objects—an intentional side effect—but has no effect on the rest of the figure. The first argument of `get_step` is the frame number.

The name of the figure and the function for updating the plot are passed to `FuncAnimation`, which calls `get_step` repeatedly to update the graph and generate the frames of the movie. We also provided additional arguments to `get_step` with the keyword argument `fargs`. These specify the data, line, and point objects to use when updating the frame. (The current frame number is automatically passed as the first argument. If the function requires additional arguments, as in this example, these should be passed by using `fargs`.)

3D animated graphs

The methods in this section also work with `Axes3D` objects, with minor changes. For example, to update frames in a 3D plot, you must replace `set_data(x,y)` with `set_data_3d(x,y,z)` to pass three-dimensional (x,y,z) data. With minor changes such as this, you can create three-dimensional animations and movies.

8.3.2 Saving animations

To create and view an animation, you don’t need any software other than Matplotlib. You can share your code with others, but they will need Python to view your beautiful work. Sometimes it is more useful to save your animation in a format that anyone can view, such as a web page or a movie file. One simple approach is to record a Python animation directly from your computer screen using the screen recording capability of your operating system.³ We will describe two additional options that give you greater control

² The keywords `lw` and `ms` are abbreviations for `linewidth` and `markersize`, respectively.

³ On macOS, use `Screenshot.app` or press <Cmd-Shift-5>. On Windows, press <Window-Alt-R>. On Linux, you can use

over the video and more options for the output file format. The first can be implemented entirely in Python. The second requires an additional piece of software called an encoder.

HTML movies

A simple approach to making an animation is to create a flip book—a series of images that create the illusion of motion when viewed in rapid succession. You may have animated stick figures in this way when you were young, or bored.

You already know how to create a series of still images using Python. You just need a way to tie them all together and display them. At press.princeton.edu/titles/32489.html, you can download a file called `html_movie.py`. This module contains a function called `movie` that creates an HTML file that will display a series of images in a continuous loop. You do not need to be online to view the animation—you just need a web browser such as Firefox, Safari, or Chrome.

The following script will create an HTML movie of two traveling waves that pass by each other.

```
# waves.py      [get code]
# Jesse M. Kinder --- 2021
""" Generate frames for an animation of moving Gaussian waves. """
import numpy as np
5 import matplotlib.pyplot as plt
# First download html_movie.py via http://press.princeton.edu/titles/11349.html
# or http://physicalmodelingwithpython.blogspot.com/
from html_movie import movie

10 # Generate waves for each frame.
# Return a Gaussian with specified center and spread using array s.
def gaussian(s, center=0.0, spread=1.0):
    return np.exp(-2 * (s - center)**2 / spread**2)

15 # All lengths are in [m], all times are in [s], and all speeds are in [m/s].
# Define the range of values to display.
x_min, x_max = -4.0, 4.0
y_min, y_max = -3.0, 3.0
# Define array of positions.
20 dx = 0.01
x = np.arange(x_min, x_max + dx, dx)

# Define the duration and number of frames for the simulation.
tmin, tmax = 0.0, 4.0
25 num_frames = 100
t = np.linspace(tmin, tmax, num_frames)

# Define the initial position and speed of Gaussian waves.
r_speed = 2.0      # Speed of right-moving wave
30 r_0 = -4.0       # Initial position of right-moving wave
l_speed = -2.0     # Speed of left-moving wave
l_0 = 4.0          # Initial position of left-moving wave
```

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```

# Generate a figure and get access to its Axes object.
35 plt.close('all')
fig = plt.figure(figsize=(6, 6))
ax = plt.axes(xlim=(x_min, x_max), ylim=(y_min, y_max))

# Create three empty line objects and grab control.
40 # The loop below will update the lines in each frame.
ax.plot([], [], 'b--', lw=1)           # Line for right-moving wave
ax.plot([], [], 'r--', lw=1)           # Line for left-moving wave
ax.plot([], [], 'g-', lw=3)            # Line for sum of waves
lines = ax.get_lines()                # Get list of 3 line objects in plot.

45 # It is essential that the frames be named in alphabetical order.
# {:03d} formats integers with three digits, including leading zeros if needed:
# '000_movie.jpg', '001_movie.jpg', and so on.
file_name = "{:03d}_movie.jpg"

50 # Generate frames and save each figure as a separate .jpg file.
for i in range(num_frames):
    r_now = r_0 + r_speed * t[i]          # Update centers of waves.
    l_now = l_0 + l_speed * t[i]
55    yR = gaussian(x, r_now)             # Get current data for waves.
    yL = -gaussian(x, l_now)
    lines[0].set_data(x, yR)              # Update right-moving wave.
    lines[1].set_data(x, yL)              # Update left-moving wave.
    lines[2].set_data(x, yR + yL)         # Update sum of waves.
    plt.savefig(file_name.format(i))       # Save current plot.

# Use HTML movie encoder adapted from scitools to create an HTML document that
# will display the frames as a movie. Open movie.html in web browser to view.
movie(input_files='*.jpg', output_file='movie.html')

```

To view the animation, open `movie.html` in a web browser. (To open a file from your computer in your web browser, choose the menu option `File>Open file ...`, and direct your browser to the file you have created.) This file must be located in the same folder as the image files to run properly. (It is not a stand-alone movie. It just tells your browser which image files to display and when to display them.) To share the animation, send the entire folder to a friend, or upload it to a web server.

In the last line, the keyword and value `input_files='*.jpg'` uses the asterisk as a wildcard to instruct the function to use *all* files in the current folder that have the `.jpg` extension. It will insert the files into the movie in alphabetical order. Line 49 formats the file names so that alphabetical and chronological order are the same. (If we had used the `{:d}` format specifier, there would be no leading zeros, and `10_movie.jpg` would come before `2_movie.jpg` in alphabetical order.)

The `html_movie` module was adapted from the `scitools.eazviz.movie` module in the `scitools` library, developed by Hans Petter Langtangen.⁴

⁴ `scitools` is a useful library, but as of this writing it is not fully compatible with Python 3. The entire library can be downloaded at github.com/hplgit/scitools. See also Langtangen, 2016.

Using an encoder

The approach in the preceding section generates an animation that you can share, but it doesn't create a standalone file in a standard video format suitable for embedding in another web page, uploading to YouTube, or editing with video software. (Perhaps you could add a soundtrack ...) To save the animation in a standard video format, you must extend Python's capabilities with a video **encoder**. FFmpeg, which can be installed using the `conda` package manager or downloaded at www.ffmpeg.org, is one Python-friendly option.

The last line of `walker.py` above, when uncommented, calls `my_movie.save`, which in turn invokes FFmpeg.

To use an encoder, you need to download and install it on your system, then make certain that Python can locate it. Using a package manager is the most straightforward way to do this. A package manager will determine all the libraries that are necessary to run the program you request, then download, install, and link everything together so your program runs properly. Section A.4 in Appendix A describes how to install FFmpeg.

Once you have successfully installed FFmpeg, you can uncomment the last line of `walker.py` above. Now when you run the script, Python will create a movie file called `random_walk.mp4` that you can edit, upload, and view in a movie player or web browser.

If you have difficulty linking Python to FFmpeg, you can run FFmpeg directly from your operating system's command line (not the IPython console). Make a series of frames with sequential names as in `waves.py` above. Then, use the following command to create the movie:

```
ffmpeg -i %03d_movie.jpg -pix_fmt yuv420p movie.mp4
```

The option `-i` option identifies the file names of the frames created by your code; the next option may be useful for movies to play on Mac computers.

ImageMagick is another free software package that can be used to create animations at the command line. The software can be downloaded at imagemagick.org, or installed by using the `conda` package manager as described in Section A.4 (page 164). If you have installed ImageMagick, you can create a `gif` animation from the output of `waves.py` by using the following command:

```
convert -delay 1x24 *.jpg movie.gif
```

The `-delay 1x24` option instructs ImageMagick to wait 1/24th of a second between each frame, and `*.jpg` tells ImageMagick to use all file names—in alphabetical order—that have a `.jpg` extension. You can view `movie.gif` in any web browser, or you can embed it in a web page.

8.3.3 Conclusion

This chapter introduced techniques for turning images into NumPy arrays, for turning arrays into images, and for creating movies with Python. In the next chapter, we will extend the idea of images as arrays to analyze experimental data.

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CHAPTER 9

Third Computer Lab

In this lab, you will import images into Python and display them. You will also explore an important operation in image analysis: convolution. If you've ever played with photographic software such as GIMP¹ (or a commercial alternative), you may have used convolutions to smooth or sharpen images but perhaps you didn't know it. Your eyes and brain are also doing convolutions whenever you look at anything.

Not all visual data comes from photographs. Many kinds of experimental measurements generate arrays of numbers associated with points in space. Such data can be communicated to our brains as images, as in electron microscopy, computed tomography, and magnetic resonance imaging. Image processing is useful for making these images more meaningful to humans.

Our goals in this lab are to

- Explore the effects of various kinds of local averaging on an image;
- See how to use such averaging to decrease noise in an image; and
- Use specialized filters to emphasize specific features in an image.

9.1 CONVOLUTION

Python gives us access to tools for manipulating images, including convolutions. However, many of the details are hidden. Before getting into Python specifics, let's take a look at the mathematical definition and properties of convolution. Convolutions arise frequently in many areas of applied mathematics, including probability, statistics, signal processing, and differential equations.² A common definition of the two-dimensional discrete convolution C of an image array I with a filter array F is

$$C_{i,j} = \sum_{k,\ell} F_{k,\ell} I_{i-k,j-\ell}. \quad (9.1)$$

The sum ranges over all values of k and ℓ that refer to valid entries of both F and I . For example, we cannot have $i = 1$ and $k = 10$ because the first index of I would be -9 . (Python may accept this, but the result would not be what you intended.)

Your
Turn
9A

- Consider the trivial transformation, for which F is a 1×1 matrix with a single entry equal to 1. Explain why C is the same as I in this case.
- Suppose that the size of F is $m \times n$, and that of I is $M \times N$. Explain why the size of C is $(M + m - 1) \times (N + n - 1)$.

¹ GIMP is freeware: www.gimp.org.

² For applications to probability distributions see Nelson, 2015, Chapter 4; for image processing, see Nelson, 2017, Chapter 11.

When we convolve an image with a filter, we get another image. The expression in Equation 9.1 is a set of instructions for constructing this new image: To create each pixel in C, we take the pixels from a subset of the original image, multiply them by their respective weights in the filter, and add up the result. It's a simple recipe that can have very different results depending on the filter.

9.1.1 Python tools for image processing

Chapter 8 explained how to load, display, and save images with PyPlot. We also performed a mathematical operation on an array to modify the image. Now we will expand our capabilities by importing a module with several image processing functions:

```
from scipy.signal import convolve
```

Use `help` to learn about this function and its options. All of the filters we will use in this lab accept similar arguments.

In this lab, you will explore several filters and convolutions. As you proceed through the exercises, you will see a photograph transformed by each operation. To get an idea of what is happening on a pixel-by-pixel level, you can apply the same convolution to a single dot. (This is the **impulse response** of the filter.) This will allow you to see the shape of the filter and better understand what you are doing to the photograph.

Try the following now:

```
from scipy.signal import convolve
impulse = np.zeros( (51, 51) )
impulse[25, 25] = 1.0
my_filter = np.ones( (3, 3) ) / 9
response = convolve(impulse, my_filter)
plt.figure()
plt.imshow(response)
```

We will explain this filtering operation in the next section. For now, compare the size of two arrays, `impulse` and `response`. They are not the same. You found in Your Turn 9A that the convolution of an image with a filter is at least as large as the original image, and usually larger. Do we want the image to increase in size? Where did the extra points come from in the mathematical derivation?

Before we answer those questions, refer back to Equation 9.1 and look at $C[0, 0]$. The allowed values of k and ℓ for this point give only one contribution. Likewise, $C[M+m-2, N+n-2]$ has only one contribution:

$$\begin{aligned} C[0, 0] &\text{ is } F[0, 0] * I[0, 0] \\ C[M+m-2, N+n-2] &\text{ is } F[m-1, n-1] * I[M-1, N-1]. \end{aligned}$$

The points at the edges of the convolved image receive contributions from fewer points in the original image than do the points in the interior. This behavior may lead to distortions at the edges of the convolved image. `convolve`'s default behavior is to return all of the points in the resulting array. But perhaps it would be better to crop the edges and return only the central portion of the convolved image. This has two advantages. First, every point in the convolved image would use at least a quarter of the filter. Second, convolution would not change the size or shape of the original image. We can get this behavior from `convolve` by supplying the keyword argument `mode='valid'`.

Even with this cropping, however, points near the edge must be treated differently from those in the center. `convolve` and its relatives offer several options. The simplest is to imagine that the image is surrounded by an infinite black border. That means we effectively enlarge our image array to whatever size it needs to be to provide the same number of points for every pixel in the convolution and set the values of all the new points to 0. This is the default behavior of `convolve`.³

9.1.2 Averaging

A very simple filter assigns the same weight to each pixel in a fixed region, as in the impulse response example above. Each pixel in the convolved image is an average of its neighbors in the original image.

Assignment:

Follow the instructions of Section 4.1 to obtain the data set `16catphoto`. Copy the files `bwCat.tif`, `gauss_filter.csv`, and `README.txt` into your working directory. Load the arrays with `np.loadtxt` and load the image into an array using `plt.imread`. (See Section 4.1.2 and Section 8.1.1.)

- a. *Make a 3×3 array `my_filter` in which each element equals $1/9$. We'll call this array the “small square filter.” Why does it make sense to choose the value $1/9$?*
 - b. *Use `convolve` to convolve your new filter with the image you downloaded, and display the result. How does the image change?*
- [Hints: You can retain the previous picture for comparison by using `plt.figure()` to create a new figure before displaying the second. Another option is to display the plots side by side with `plt.subplots`. (Consult `help(plt.subplots)` and Section 4.3.9.) You must set the color map separately for each figure that you create. (Consult `help(plt.colormaps)`.)]*
- c. *Repeat part (a) using a 15×15 array (the “large square filter”). Be sure to use a constant value appropriate to this larger array. How does the image change? How does it compare with the result of the smaller filter?*
 - d. *Use the definition in Equation 9.1 to derive an expression for the value of a specific pixel—say $(100,100)$ —in the convolved image, in terms of the elements of the small square filter and the original image array. Show that it is the average of some of the neighboring pixels in the original.*

9.1.3 Smoothing with a Gaussian

Now we will look at a more complex filter, called a Gaussian filter. Load `gauss_filter.csv` into a NumPy array called `gauss`. Then, review Section 6.4.3 to refresh your memory on the function `plot_surface`.

Assignment:

- a. *Display the convolution of `gauss` with the original image.*

³  For more options, use `scipy.signal.convolve2d` instead. You can control the treatment of edges with the `boundary` and `fillvalue` keyword arguments.

- b. Use `plt.imshow` to compare the convolutions of a single dot with a Gaussian filter and with the square filter you used in question 9.1.2b. The `impulse` filter above is an image of a single dot against a black background.
- c. Use `plot_surface` to view the convolved images from part (b) in three dimensions. This is actually a plot of the filters themselves. Use the definition of convolution to explain why. Then, explain how convolution with a Gaussian filter differs from a square filter. When might you want to use a Gaussian filter instead of square filter?

9.2 DENOISING AN IMAGE

Measuring instruments, including your eyes, inevitably introduce some randomness, or “noise.” You can simulate this effect by making a noisy version of the original image you imported. To do this, multiply each pixel in the original image by a random number.

Assignment:

- a. Multiply each pixel of the original image by a random number between 0 and 1. Compare this noisy image with the original.
- b. Apply each of the three filters from Sections 9.1.2–9.1.3 (small square, big square, Gaussian) to the noisy image from part (a). Do they improve the image? If so, which one works the best? Why? Zoom in on a small region of the resulting images. How do they compare at this scale?

9.3 EMPHASIZING FEATURES

You have probably heard news people say, “Geeks at NASA’s Jet Propulsion Laboratory have enhanced these images....” Let’s join the fun.

In between “features” (real things of interest to us) and “noise” (random things), experimental images may contain things that are real, but not of interest to us. We may wish to de-emphasize such things, or we may wish to quantify some visual feature. A. Zemel and coauthors encountered such a situation when making fluorescence images of mesenchymal stem cells. When subjected to mechanical stress (stretching), the cells polarize: The internal network of “stress fibers” begins to align in the direction of the stretch.⁴ Zemel and coauthors sought to quantify the extent to which the cell was polarized, at every point in the cell.

Follow the instructions of Section 4.1 (page 53) to obtain the data set `17stressFibers`. Copy `README.txt` and `stressFibers.csv` into your working directory.

Load and plot the data. The image shows the stress fibers. We will now construct and apply a filter that emphasizes long, slender objects that are oriented vertically.

Assignment:

- a. Execute the following code, then make a surface plot of the filter. Describe its significant features.

⁴ A fluorescent label for nonmuscle myosin IIa was used to tag the stress fibers. See Zemel et al., 2010.

```
# convolution.py      [get code]
v = np.arange(-25, 26)
X, Y = np.meshgrid(v, v)
gauss_filter = np.exp(-0.5*(X**2/2 + Y**2/45))
```

- b. Use the following “black box” code to modify your filter from part (a), then make a surface plot of the resulting filter. Compare and contrast `combined_filter` and `gauss_filter`.

```
laplace_filter = np.array( [ [0, -1, 0], [-1, 4, -1], [0, -1, 0] ] )
combined_filter = convolve(gauss_filter, laplace_filter, mode='valid')
```

The matrix `gauss_filter` emphasizes features that are long, slender, and oriented vertically. Other features are averaged out. The array `combined_filter` accentuates the edges of such objects.⁵

- c. Now use `convolve` to apply the filter to the fiber image, display your results, and comment.

You may notice that the filtered image has poor contrast. After convolution, the values assigned to some pixels are extremely large or extremely negative, but most points fall in a narrow range between the extremes. Python uses its shades of gray to interpolate between these extremes, giving most points a gray level somewhere in the middle of the range. You can highlight the features you wish to emphasize with another modification.⁶

```
plt.imshow(image, vmin=0, vmax=0.5*image.max())
```

- d. To emphasize **horizontal** objects, repeat the above steps with a different choice for `gauss_filter`.
Optional: Make two more filters that emphasize objects oriented at $\pm 45^\circ$ with respect to vertical.

9.4 T2 IMAGE FILES AND ARRAYS

When we load an image to an array, Python does not know the array represents an image. It will apply the same rules of arithmetic to an “image array” as to any other, including changing its data type. For example, if you multiply an array of unsigned integers (`uint8`) by an array of floats (`float64`), NumPy returns an array of floats. For plotting in Python, this is usually not a problem. However, if you plan to process the transformed image in some other application, it may be.

You need a `uint8` array for some image processing operations. Moreover, the data type of an array controls how the array is saved to an image file. A `float64` array gets saved to a 4-layer TIFF, whereas a `uint8` array gets saved to a 1-layer grayscale TIFF by the Pillow library. A black-and-white image could inadvertently be converted to full color!

We saw an example of this in Section 9.2. The default return value of `np.random.random` is a random floating point number with mean 0.5. If you multiply an image by a random array of this type, three things will happen: the resulting image will get darker, its maximum value will be less than 255,

⁵ T2A Laplace filter emphasizes edges, but is sensitive to noise. A Gaussian filter smooths out noise and edges. Combining the two creates a *Laplace of Gaussian* (LoG) filter that emphasizes edges while suppressing noise. The “elongated” Gaussian used here creates an eLoG filter that emphasizes the edges of objects with a particular alignment.

⁶ T2The keyword arguments implement a “window/level transformation.” Pixels whose luminance is less than `vmin` are displayed in black, those whose luminance is greater than `vmax` are displayed in white, and pixels whose luminance lies between these extremes are displayed in shades of gray. This transformation emphasizes contrast in the range of interest while ignoring features

outside that range.

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and the resulting array will contain floats instead of integers. For some operations, like convolution with an eLoG filter, some of the elements of the array may even be negative. An array like this may cause problems in applications that expect arrays like those returned by `plt.imread`—arrays whose entries are integers between 0 and 255.

We can transform any array to the proper image format with three steps:

1. Shift the minimum of the array to zero. That is, find the smallest value of the array, and subtract that value from every element.
2. Rescale the array so that the maximum value is 255.
3. Change the data type to `uint8` using the `astype` method of the array: `a = a.astype('uint8')`.

The `scipy.ndimage` library contains many functions for working with images, including convolutions and filters that preserve the data type of the image. If your image files need to remain `uint8` image files, explore this library. Be sure to consult the documentation, though. The default behavior of `scipy.ndimage.convolve` differs from `scipy.signal.convolve`.

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CHAPTER 10

Advanced Techniques

*We can only see a short distance ahead,
but we can see plenty there that needs to be done.*
— Alan Turing

This chapter introduces sophisticated tools and techniques for physical modeling. It can be skipped on a first reading. You can already do a lot with what you have learned!

To illustrate our points, we will develop a case study involving “first passage” problems, which arise throughout physics, chemistry, and even finance. First, we describe Python data structures and syntax that can help simplify and organize large programs and simulations. Next, we introduce Python tools for data science. Then, we introduce computer algebra: working with symbols instead of numbers. We close by showing you how to write your own classes with Python—and why.

10.1 DICTIONARIES AND GENERATORS

Data structures organize the data in a program. There is not one data structure that is superior to all others; each has its strengths and weaknesses. We have already looked at some of Python’s built-in data structures: lists, strings, and tuples. But we focused primarily on NumPy arrays, since these are the cornerstone of much scientific computing. Numerical arrays make efficient use of memory for large data sets, and they allow fast, parallel calculations on their elements. However, they are not as efficient as lists when inserting and removing elements, and a numerical array is ill-suited to working with text. We use strings for this. Choosing the right data structure for a task can simplify your code, improve its performance, and reduce errors.

In this section, we will discuss one more useful data structure: dictionaries. Next, we introduce a useful shortcut for creating lists. We close this section by introducing convenient ways to pass collections of arguments to functions.

First passage

Suppose that you wish to know how long it takes for a particle undergoing a one-dimensional random walk to reach a point 100 steps to the right of its initial position.¹ This is an unlikely event, unless we take a lot of steps. (After 100 steps, the odds are just 1 in 2^{100} .) We don’t know offhand how many total steps are required, or indeed whether some particles might never reach this point. We can instead limit ourselves to a large but finite number N of steps, and then determine when—if ever—a particle first passes the point $x = 100$ or some other target distance L . For generality, we also introduce a third parameter to allow us to explore biased random walks.

¹ Section 6.2.2 (page 82) and Chapter 7 introduced random walks.

```

# first_passage.py      [get code]
def first_passage(N, L, p=0.5, message=False):
    """
    Return number of steps for first passage of x==L,
    or give up after N and return np.nan.

    The walker takes steps to the right with probability p.

    Use message=True to display the results.
    """
    rng = np.random.default_rng()      # create a random number generator object
    dx = 2*(rng.random(N) < p) - 1   # individual steps
    x = np.cumsum(dx)                 # location after each step
    at_target = np.nonzero(x==L)[0]   # indices where x == L

    if at_target.size > 0:
        n = at_target[0] + 1
        if message:
            print("First passage of x={} occurred after {} steps.".format(L, n))
        return n
    else:
        if message:
            print("Did not reach x={} after {} steps.".format(L, N))
        return np.nan

```

(We are going to be putting this function inside large `for` loops later, so the `print` commands are disabled by default. Pass the keyword argument `message=True` to display a message about the result.) Calling `first_passage(10**6, 100, message=True)` a few times reveals that sometimes the walker fails to reach the target even after one million steps, but most of the time it does so in far fewer steps.

We will build on this simulation as we illustrate the use of dictionaries, generators, data science libraries, and Python classes.

10.1.1 Dictionaries

With an array or a list, we look up elements using an index. We provide Python with an integer, and it uses this integer to report what is stored at a particular location in an array. For example, `x[3]` returns the fourth² element of the array or list called `x`. This scheme is convenient for ordered numerical data, like the values of a function over a range of inputs. However, sometimes it is more convenient to store data as **key-value pairs**. For this purpose, a **dictionary** is the appropriate data structure.

Suppose that we wish to store the input parameters for our first passage problem. One approach would be to assign each parameter to a variable:

```

N = 1000
L = 10
p = 0.5

```

² Recall that Python's indexing is based on the idea of offsets (Section 2.2.5, page 25). The first element in the array is `x[0]`, so

`x[3]` is the third item *after* `x[0]`, that is, the fourth element in the array.

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Calling `first_passage(N,L,p)` will run the simulation with these parameter values. This simple approach can become cumbersome if you wish to run multiple simulations with different parameters or many simulations with the same parameters: Eventually you have lots of results and must keep track of which results came from which set of parameters.

You could keep a record of your work in a lab notebook. A more elegant approach is to create a Python dictionary to store the parameters:

```
parametersA = { 'N': 1000, 'L': 10, 'p': 0.5 }
```

`parametersA` stores its data as pairs: a **key**, used to look up the element, and a **value** associated with that key. We can look up a value by providing its key:

```
print(parametersA['N'])
print(parametersA['p'])
```

A dictionary is mutable, so you can change its values, too:

```
print(parametersA['L'])
parametersA['L'] = 20
print(parametersA['L'])
```

This example shows the simplest way of constructing a dictionary: a series of `key:value` pairs, separated by commas, enclosed in curly brackets. You can use any immutable object for a key: a string, an integer, a float, a tuple. Any Python object, immutable or not, can be stored as a value. You look up an element by providing a key as an index in square brackets—just like providing an index to an array.

An alternate method of constructing a dictionary resembles keyword arguments in functions.³

```
parametersB = dict( N=1000, L=10, p=0.5 )
```

This second method is convenient because the construction of the dictionary parallels a function call. The connection is not superficial: As we will see shortly, a dictionary can later be passed to a function and unpacked as a set of keyword arguments.

There are three common ways to iterate over the contents of a dictionary.

```
# Iterate over keys.
for k in parametersA.keys(): print( "{} = {}".format(k, parametersA[k]) )

# Iterate over values.
for v in parametersA.values(): print( "{} squared is {}".format(v, v**2) )

# Iterate over key-value pairs.
for k,v in parametersA.items(): print( "The value of {} is {}".format(k,v) )
```

The order of the printed keys and values may be different from the order you used to create the dictionary.

When should you use a dictionary? If you have an unordered data set that can be represented as a collection of keys and values, consider using a dictionary instead of a list or an array. In particular:

- When you inspect a dictionary, for example with `print`, seeing the key and the value together can help you understand what you see.
- Dictionaries allow fast insertion, removal, and searching, even when the number of entries is large.⁴

³ See Section 6.1.3

⁴  Data is stored in an unordered hash table. Lookup, insertion, and deletion are $\mathcal{O}(1)$, independent of the dictionary's size.

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- Dictionaries can be an efficient way to store large sparse matrices, too. If you have a large array and most of the values are zeros, such as an adjacency matrix for a very large graph, using the nonzero indices as keys and the elements as values can save a lot of memory and searching time.
- Dictionaries are convenient for storing the parameters of a simulation, as we will now illustrate.

You can use dictionaries to organize data within a script, or even at the IPython command prompt. Try the following:

```

data = {}
data['A'] = { 'input': parametersA,
             'results': first_passage(parametersA['N'],
                                         parametersA['L'],
                                         parametersA['P']) }
5
data['B'] = { 'input': parametersB,
             'results': first_passage(parametersB['N'],
                                         parametersB['L'],
                                         parametersB['P']) }
10 print(data)

```

In this example, `data` is a dictionary whose two entries are themselves dictionaries, each with two subentries.⁵ Note that in contrast to lists or arrays, the target of an assignment may be an entry whose key does not yet exist in the dictionary. You can *assign* nonexistent elements of a dictionary, but you will get a `KeyError` if you ask for one. Type `data['C']` and press <Return> to see this kind of error.

We now have two complete simulations stored in a single, easily accessible object. We can check the input parameters for a given result, if needed, or we can add more simulations with a particular set of input parameters if we find an interesting result.

The next two sections will introduce Python methods that can save typing and lead to more reproducible code, as we start to generate even more data for our example problem.

10.1.2 Special function arguments

Starred expressions

In earlier chapters, we have called `plt.plot` with several different sets of arguments, yet each time we got a graph with the options we intended. Consulting Python's built-in help, or its online resources, we also notice some strange-looking arguments. For example, the output from `help(plt.plot)` shows a generic argument list that starts with `*args`. In help text, `*args` means "any number of positional arguments." Thus, `plt.plot(x,np.sin(x))` and `plt.plot(x,np.sin(x),'r-',x,np.cos(x),'g--')` both work fine, and we can add many more elements if we would like. We will return later to the last item mentioned in the help text, `**kwargs`. First let's take a closer look at `*args`.

You can use the star syntax yourself: If you create a list called `myargs`, you can pass its *individual elements*—not the list itself—as arguments to a function as `*myargs`.⁶ For example, you can organize your functions and formatting before calling the plot command:

```

t = np.linspace(-2*np.pi, 2*np.pi, 201)
line1 = [t, np.sin(t), 'r-']    # Store domain, function, and format in a list.
line2 = [t, np.cos(t), 'k--']

```

⁵ One of *those* subentries is also a dictionary, with three subsubentries!

⁶ You can also use a tuple, array, or string.


```
plt.plot(*line1, *line2)
```

We placed each set of plot arguments into a list, and then used a **starred expression** to pass them to `plt.plot`. This syntax also allows you to use the same parameters in multiple plot commands without explicitly typing them in multiple locations, and hence to maintain uniform style. (*Define once; reuse often.*) In short:

A starred argument means “use each element of this object as a separate positional argument.”

You can use starred expressions in any function with positional arguments. Let’s use this syntax to call `first_passage` several times:

```
parametersC = (1000, 25, 0.5, True)
for i in range(20): first_passage(*parametersC)
print(parametersC)
```

(The fourth element of `parametersC` instructs `first_passage` to print the result to the screen.)

Double-starred expressions

Starred expressions are convenient, but the example just given is a bit cryptic. It is not explicit about the meaning of the input parameters. Another kind of special argument, the **double-starred expression**, allows us to pass keyword arguments via a dictionary.

In Python documentation, `**kwargs` means “any number of keyword/value pairs—or none at all.” We have used this option in many plot commands already:

```
plt.plot(x, np.sin(x), linewidth=4, label='Sine Wave')
```

The documentation of `plt.plot` mentions `linewidth`, `label`, and many other keywords that do not appear in the arguments list of `plt.plot`. These are all included in `**kwargs`.

You can use the double-star syntax to pass a collection of key/value pairs to a function via a dictionary.

```
parametersD = dict(N=1000, L=25, p=0.5, message=True)
for i in range(20): first_passage(**parametersD)
print(parametersD)
```

This approach saves typing every time we call `first_passage`, as did the use of `*args` above. As mentioned previously, `print(parametersD)` is also much more informative than `print(parametersC)`. In short,

A double-starred argument means “use each entry of this dictionary as a separate keyword argument.”

You can write your own functions that accept variable numbers of positional and keyword arguments by using the `*args` and `**kwargs` syntax. You can learn more about this and other subtleties of function arguments at docs.python.org/3/tutorial/controlflow.html.

10.1.3 List comprehensions and generators

While running the sample codes up to this point, you may have noticed some trends and surprises in how long it takes the random walker to reach its target and how often it fails to reach the target. Our next step in the analysis is to build up a data set and try to get a more quantitative understanding of these trends.

One approach is to use a `for` loop to populate a list of results in our data dictionary.

```
samples = 100
data['A']['results'] = []    # Create empty lists to store the data.
data['B']['results'] = []
for i in range(samples):
    # Append new simulation results to the existing list.
    data['A']['results'] += [first_passage(**data['A']['input'])]
    data['B']['results'] += [first_passage(**data['B']['input'])]
print(data['A']['results'])
print(data['B']['results'])
```

The reason for using a list is that we may want to extend the data set. Running the loop again (but not the lines above it that would clear the results!) will add 100 more points to the data set.

List comprehensions

There is another way to create a Python list:

```
squares = [n**2 for n in range(100)]
```

This construction is called a **list comprehension**. List comprehensions are not essential to Python programming—you can accomplish the same thing with a `for` loop. But this syntax is clear and concise:

A list comprehension is a one-line `for` loop enclosed in square brackets.

Returning to our case study, we can use list comprehensions to organize our entire simulation in a few lines. Starting from the top, we have

```
# data_dictionary.py      [get code]
data = {}          # empty dictionary to store all data
data['A'] = {}    # empty dictionary within data to store Simulation A
data['B'] = {}    # empty dictionary within data to store Simulation B
5
# Define and run simulations.
samples = 500

data['A']['input'] = dict(N=1000, L=10, p=0.5)
10 data['A']['results'] = \
    [ first_passage(**data['A']['input']) for n in range(samples) ]

data['B']['input'] = dict(N=1000, L=20, p=0.5)
data['B']['results'] = \
15   [ first_passage(**data['B']['input']) for n in range(samples) ]

# Run more simulations.  Use "+=" to append new list to old.
data['A']['results'] += \
    [ first_passage(**data['A']['input']) for n in range(samples) ]
20 data['B']['results'] += \
    [ first_passage(**data['B']['input']) for n in range(samples) ]
```

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Turn
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We now have a start on a data set for investigating first passage. Add more simulations with different parameters. Generate walks with more steps and different target distances. Store them all in the same dictionary.

Generators

List comprehensions illustrate a broader class of Python objects that can be useful in managing memory in your programs.

Looking at an expression like `x = [n**2 for n in range(20)]`, you might ask yourself, “What exactly *is* the thing inside the brackets?” Its meaning may be clear enough to a human reader, but it is not a list, a string, or an array. Inside the definition of a list, “`n**2 for n in range(20)`” is called a *generator expression*. It specifies an object called a **generator**. Like many generators, this one is constructed using `range`. `range` is not a generator, a list, or an array. It creates yet another type of Python object that can produce a sequence of integers.⁷ You can think of a generator or a `range` object as specifying a *rule* for generating a sequence rather than storing the actual elements of that sequence.

You can create a generator outside of a list, too:

```
G = ( n**2 for n in range(100) )
print( type(G) )
```

The `print` command reveals that `G` is not a tuple, despite the round parentheses. Unlike a tuple, a list, or an array, a generator cannot be indexed: `G[0]` will raise an exception. About all a generator can do is return the *next* element of its sequence, following its rule:

```
print(next(G))
print(next(G))
print(next(G))
```

When you run these commands, you will see that every time we call Python’s built-in `next` function on `G`, we get the next term in the sequence of squares. We can keep calling `next(G)` until we get to the end of its sequence—in this case, the integer $99^2 = 9801$. After this, calling `next(G)` will raise a **StopIteration** exception.

Why use generators? Their utility lies in their compactness. Generators embody a concept in computer science known as *lazy evaluation*. Rather than compute the entire sequence and store it, Python computes elements of the sequence only as they are needed (and thus only *if* they are needed). Computer memory is cheap, but if the storage required for the objects in your calculation exceeds your computer’s available RAM, you will notice a dramatic increase in the time it takes to finish a calculation.

Suppose that you need a million random numbers. You could store all of these numbers in an array or a list, or you could use a generator to produce them as needed. The following example uses the built-in `__sizeof__()` method to display the size of each object, in bytes:

```
N = 10**6                      # number of data points
rng = np.random.default_rng()    # create a random number generator object
```

⁷ In Python 3, `range` creates a `range` object. In Python 2.7, `range` creates a list of integers. The behavior of `xrange` in Python

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```
r_array = rng.random(N)                      # Store in NumPy array.
5 r_list  = [ rng.random() for n in range(N) ] # Store in Python list.
r_iter  = ( rng.random() for n in range(N) )  # Store in generator.

print( "Size of array: {}".format( r_array.__sizeof__() ) )
print( "Size of list: {}".format( r_list.__sizeof__() ) )
10 print( "Size of generator: {}".format( r_iter.__sizeof__() ) )
```

We see that the NumPy array and the Python list each use over 8 MB to store the data, while the generator only requires 96 bytes! The savings come at a cost, though. You cannot vectorize code that uses generators, nor can you access an arbitrary element using an index. However, if you only need a sequence of values one at a time, you can free up memory by using generators. This approach is also useful if you won't necessarily need all of the elements of the sequence.

Don't create a very large array or list just to iterate over it.

Use `range` or a generator.

The criterion for “very large” will change as technology evolves, but the principle will still apply.⁸

Enumeration

Arrays, strings, lists, tuples, dictionaries, range objects, generators, and many other Python objects can all be scanned in a `for` loop: `for item in object:` Any such object is called an **iterable**, because you can “iterate” its elements (access them in sequence). Some iterables do not have a natural order among their elements: for example, the keys of a dictionary. Nevertheless, it is often useful to count the elements of a collection as you access them—to **enumerate** them. Python provides a built-in function called `enumerate` to do exactly this.

```
L = [ n**2 for n in range(10) ]
for x in enumerate(L): print(x)
```

From this example, we see that `enumerate` accepts an iterable (here the list `L`) and returns a new iterable consisting of a tuple for every item in `L`. The first item of each tuple is an integer counter, and the second element is the corresponding item from `L`. By unpacking this tuple, we can separate the counter from the element and use them independently. The behavior of `enumerate` is the same for any type of iterable.

Enumeration can be useful when you want to link the elements in one collection with those in another: dictionary keys and columns of an array, lines of a file and elements of a list, functions and subplots. The following example uses the keys of a dictionary as labels for functions in a plot and titles of subplots:

```
theta = np.linspace(-2*np.pi, 2*np.pi, 201)
functions = { r"$\sin \theta$": np.sin(theta),
              r"$\sin^2 \theta$": np.sin(theta)**2,
              r"$\cos \theta$": np.cos(theta),
5             r"$\cos^2 \theta$": np.cos(theta)**2 }
styles = ['r-', 'g--', 'b:', 'k-.']

# Plot functions in same figure, with labels.
plt.figure()
```

⁸ Running Python on a laptop with 8 GB of RAM, “very large” is a NumPy array of about 10 million elements.

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```
10 for n, k in enumerate(functions.keys()):
    plt.plot(theta, functions[k], styles[n], label=k)
plt.legend()

# Plot functions in separate subplots, with titles.
15 fig, ax = plt.subplots(2, 2, sharex=True, sharey=True)
for n, k in enumerate(functions.keys()):
    I, J = n // 2, n % 2      # Use modular arithmetic to get subplot indices.
    ax[I, J].plot(theta, functions[k])
    ax[I, J].set_title(k)
```

NumPy provides a related function called `np.ndenumerate` that generates the indices and elements of an `ndarray`, as well as `np.ndindex`, which only generates the indices for a given shape.

```
the_shape = (4, 4)
R = np.random.random(the_shape)
for I, r in np.ndenumerate(R):
    print("The element at {} is {:.3f}.".format(I, r))

5
R1 = np.random.random(the_shape)
R2 = np.random.random(the_shape)
for I in np.ndindex(the_shape):
    print("The elements at {} are {:.3f} and {:.3f}.".format(I, R1[I], R2[I]))
```

`enumerate` and its relatives can be extremely useful when iterating over several objects simultaneously.

10.2 TOOLS FOR DATA SCIENCE

Python is an extremely popular platform for data science, machine learning, and deep learning. While a full discussion of these topics is beyond the scope of this book, let’s take a quick look at the basic tools: the `pandas` and `scikit-learn` libraries, contained in modules called `pandas` and `sklearn`.⁹ We will apply these tools as we continue to develop our first passage case study.

10.2.1 Series and data frames with pandas

A NumPy array is a very useful data structure for working with numerical data in the form of arrays and matrices. However, this structure is rigid: All of its elements must have the same data type, and it provides no way of labeling the data within. If you import data from a spreadsheet, for example, you have to keep track of column labels elsewhere, perhaps via a separate `readme.txt` file. Even importing the values can be a challenge. If there are empty columns in a spreadsheet, you will likely end up with errors when you import it using `np.loadtxt` (Section 4.1.2). All of these challenges are part of the daily workflow in data science, and so a special set of tools has been developed to address them in the Python ecosystem. Most of these tools are part of the `pandas` library. We will only look at the most basic elements of `pandas`.

⁹ If you did a full installation of Anaconda, you should have these libraries already. (See Appendix A.) If not, run “`conda install pandas scikit-learn seaborn`” at the command line before attempting the exercises in this section.

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First, let’s transform our list of data points into a NumPy array and a pandas Series object to see some of the similarities and differences:

```
import pandas as pd

dataArray = np.array( data['A']['results'] )
dataSeries = pd.Series( data['A']['results'] )

print(dataArray)
print(dataSeries)
```

You will see slightly different output, but the two objects we created seem to be quite similar at first glance. They are. From `help(pd.Series)`, we learn that a series is a “one-dimensional ndarray with axis labels.” Indeed, you can do everything with a series that you could do with a NumPy array.

Upon closer inspection, however, **Series** objects have additional methods, and slightly different behavior for methods of the same name. As an example, let’s look at statistics for our random walks:

```
print("Average steps to reach x={} ...".format(data['A']['input']['L']))
print("ndarray: ", dataArray.mean())
print("Series: ", dataSeries.mean())
```

We already see a difference in behavior here. `first_passage` returns `np.nan` when the walker fails to reach its target. This indicates an invalid data point. NumPy does not want these values “swept under the rug,” because they often indicate a numerical calculation gone wrong. Thus, it propagates them through calculations: `np.nan+1` yields `np.nan`, `2*np.nan` yields `np.nan`, and so forth. When calculating the mean number of steps, the ndarray therefore returns `np.nan`. In contrast, the pandas **Series** returns a numerical value. That’s because data scientists often encounter missing data or poorly formatted data sets. Rather than declare calculations on such data sets a lost cause, the default behavior in pandas is to ignore `np.nan` and drop these entries from the calculation. This can make life easier, but it can also bias a data set, if the `np.nan` values meant something other than a missing piece of data—as is the case in our problem. Still, knowing the average number of steps to reach $x = 10$ among walkers that did so in fewer than 1000 steps is better than knowing nothing at all.

Computing statistics on a column of numbers is a very common task in data science, and so a pandas series object has a convenience function to compute them: `dataSeries.describe()` will print out a short report on the data within. Series objects also have their own plotting functions: `dataSeries.hist()` will plot a histogram of the data; `dataSeries.plot(kind='density')` will estimate and plot an estimated probability density function for the data. You can use these methods to investigate the first passage data you have collected so far.

A **Series** is a very useful data structure, but the workhorse of Python data science is the data frame. A **DataFrame** is a two-dimensional table of data, similar to a spreadsheet. You can access rows or columns, apply functions to ranges of data, search, sort, plot, and more. Here again there are similarities to a two-dimensional NumPy array. However, you can mix data types and access data by row and column labels or integer indices. Let’s construct a **DataFrame** to get more familiar with this data structure.

There are many ways to construct a `Dataframe`. One of the simplest is to provide a dictionary whose keys are the column labels and whose values are the elements of the column.

```
df = pd.DataFrame({ 'L=10':data['A']['results'], 'L=20':data['B']['results'] })
print(df.head())
```

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In the last line, `head()` is a method of `Series` and `DataFrame` objects that will extract the first few entries. (Likewise, `tail()` extracts the *last* few.) It is a useful way to inspect the contents of a data frame, and a good check on whether creating or importing a data frame was successful. You can add more columns, as you would add new elements to a dictionary:

```
df['L=30'] = pd.Series([first_passage(N=1000,L=30,p=0.5) for n in range(2000)])
```

Try some of the analysis functions on the data frame:

```
df.describe()
df.hist()
df.plot(kind='density')
```

We get a separate analysis of each column of the data set.

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Explain in what sense the statistics get worse as the target distance increases. To address this problem, repeat the construction of the data set, but with more steps and more data points: `N=10**6` and `samples=10**5`. Create histograms and probability density plots. Describe trends you see in the data for increasing L .

Pandas can read and write series and data frames to a variety of formats, but the comma-separated value file (`.csv`) is the standard choice in data science. To save your data set in this format, use the `to_csv()` method; to load, use the `read_csv` function:

```
df.to_csv("first_passage.csv", index=False)
backup = pd.read_csv("first_passage.csv")
```

You can verify that `df` and `backup` contain the same data by inspection, plotting, or statistical analysis. (A direct test for equality like `df==backup` will not work because `np.nan==np.nan` evaluates to `False`.)

We can only scratch the surface of pandas here, but you can already start using `Series` and `DataFrame` objects in your work. These objects are quite efficient in working with very large data sets, and they combine many useful features of spreadsheets, databases, and numerical arrays. The `help()` documentation is thorough and contains many useful examples. Other places to learn more include the “10 Minutes to pandas” tutorial at pandas.pydata.org; VanderPlas, 2017; and McKinney, 2018.

10.2.2 Machine learning with scikit-learn

When analyzing experimental or numerical data, we often wish to fit a model to the data. We do this by adjusting parameters in the model so that its predictions match the data as closely as possible. (You did this by hand in Chapter 5.) Once we have adjusted the parameters of the model, we can use it to predict the values of new measurements or classify new measurements. For example, we might use a series of measurements of the position of a comet to create a model of its path around the Sun. We could then use this model to predict when the comet will return. Or, we might measure the conductivity of a material at different temperatures and pressures and use a model to construct a phase diagram—to predict whether other temperatures and pressures will result in superconductivity.

Using data to make predictions is very general and extends well beyond the physical sciences. A set of mathematical tools has been developed for the process, called **machine learning**. The pandas library is designed to read, write, and analyze data. Another library, called **scikit-learn**, is designed for machine learning. (We will call it by its module's name, `sklearn`.) The two libraries work well together. In this section, we will demonstrate linear regression to study our first-passage data. However, many of the models in `sklearn` use the same protocol, so you can adapt these examples to explore other models for the same data.

Linear regression is the process of fitting a line to a data set, familiar to all lab students. Sometimes we fit a line with a ruler; more often, we use software to do the fitting for us. Let's see how this process works in `sklearn`. We will fit three linear models to our first-passage data to better understand the probability of first reaching $x = L$ after N steps.

The first step in the process is to import the model we wish to use and create an instance of it. Linear regression is part of the `linear_model` module within `sklearn`.

```
from sklearn.linear_model import LinearRegression
model = LinearRegression()
```

The next step is to prepare our data. Models from `sklearn` are particular about the shape of data sets, but they are consistent. Each data point is assumed to be of the form $(x_1, x_2, \dots, x_N, y_1, y_2, \dots, y_M)$. The $\{x_i\}$ are the inputs (independent variables)—often called “features” in the current language of data science. The $\{y_i\}$ are the outputs (dependent variables), often called “targets.” The goal of a model is to predict targets from a new set of features, based on “training data” (a collection of known feature/target data).

A linear regression model makes its predictions by using the method of least squares. To pass our training data to a model, we must break it into two arrays: `X`, an array where each row is a set of features in a single measurement, and `Y`, an array where each row is the corresponding set of targets.

For the first-passage problem, we can bin the data and try to fit the number of counts in each bin to its midpoint. This will give us a model for the probability of taking exactly N steps to first reach $x = L$.

```
steps = df['L=10'].dropna()                      # Discard walks that did not reach L.
counts, edges = np.histogram(steps, bins=20)
centers = 0.5*(edges[:-1] + edges[1:])  # Use centers of bins as features.

# Prepare data for model.
X = centers.reshape(-1, 1)                         # Reshape 1D array to column vector.
Y = counts
```

It is essential to reshape the array of bin locations into a column vector. (See Section 2.2.9.) Now we are ready to fit the model to the data:

```
model.fit(X, Y)
```

Nothing appears to have happened, but the model has “trained” itself by adjusting its internal parameters. You can access those parameters as data fields:

```
print(model.coef_)
print(model.intercept_)
```

The slope of the line is stored in `model.coef_` and the intercept is stored in `model.intercept_`. (The underscore after the name is a convention that means you should not modify these data fields.) We

can ask the model to assess its fit to the data:

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```
model.score(X, Y)
```

This returns the “coefficient of determination,” sometimes called the “ R^2 value.” A value of +1 indicates a perfect linear fit, and values near 0 or below indicate a poor linear fit. The fit is poor here, because the relationship is not linear. A graph makes this apparent:

```
xFit = np.linspace(X.min(), X.max(), 201).reshape(-1, 1)
yFit = model.predict(xFit)
plt.figure()
plt.plot(X, Y, 'ro', label="Data")
5 plt.plot(xFit, yFit, 'k-', label="Fit")
plt.legend()
```

This example shows how to use the model to predict values for new points: You pass an array of points to the model’s `predict` method.

We can explore many mathematical relationships with linear models.¹⁰ If $y = Ae^x$, then we expect a linear relationship between $\ln y$ and x . If $y = Ax^b$, then we expect a linear relation between $\ln y$ and $\ln x$. We can quickly explore these relationships:

```
# Bin data again, over smaller range.
steps = df['L=10'].dropna()                      # Discard walks that did not reach L.
counts, mybins = np.histogram(steps, bins=20)
centers = 0.5*(mybins[:-1] + mybins[1:])  # Use centers of bins as features.
5
# Don't take logarithms of zero. Exclude invalid points from all models.
valid = (centers>0) * (counts>0)
X = centers[valid].reshape(-1, 1)
Y = counts[valid]
10 logX = np.log(X)
logY = np.log(Y)

xFit = np.linspace(X.min(), X.max(), 201).reshape(-1, 1)

15 # Plot data.
plt.figure()
plt.plot(X, Y, 'ko', label="Data")

# Check for linear relationship.
20 model.fit(X, Y)
print("R2 Linear: ", model.score(X, Y) )
yFit = model.predict(xFit)
plt.plot(xFit, yFit, 'r-', label="Linear Model Fit")

25 # Check for exponential relationship.
model.fit(X, logY)
print("R2 Exponential: ", model.score(X, logY) )
yFit = model.predict(xFit)
plt.plot(xFit, np.exp(yFit), 'g-', label="Exponential Model Fit")
```

¹⁰ See Section 4.3.2 (page 63).

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```

30 # Check for power-law relationship.
model.fit(logX, logY)
print( "R2 Power Law: ", model.score(logX, logY) )
yFit = model.predict(np.log(xFit))
35 plt.plot(xFit, np.exp(yFit), 'b-', label="Power Law Model Fit")

plt.legend()

```

Your Turn
10C

Try it! Gather data and fit models to $\langle N \rangle$ versus L . That is, let X be the target location L and Y the average number of steps $\langle N \rangle$ to reach the target location. Do your data suggest a linear relationship, an exponential relationship, a power law, or something else?

10.2.3 Next steps

This is only a taste of what you can do with pandas and scikit-learn. You now know how to read and write a large number of data file types, to create models from data, and to start exploring data science with Python. You can learn a lot more from these tutorials:

- **pandas:** Searching and merging large data sets are very useful tools to add to your repertoire:
pandas.pydata.org/docs/getting_started/intro_tutorials
- **scikit-learn:** Many models follow the same pattern as the [LinearRegression](#) example in this chapter. You can apply them simply by importing them and substituting them for [LinearRegression](#):
scikit-learn.org/stable/tutorial/index.html
- **seaborn:** This is a popular plotting library in the Python data science world. It works well with pandas and scikit-learn:
seaborn.pydata.org/tutorial.html
- **TensorFlow and Keras:** If you are interested in deep learning with Python, these are the libraries to learn. TensorFlow is a library of machine learning tools, and Keras is a high-level interface for working with those tools:
tensorflow.org/tutorials

10.3 SYMBOLIC COMPUTING

Physical modeling often requires symbolic analysis—not just crunching numbers. You need to derive formulas and solve equations using symbols instead of numbers. This is often done with a pencil and paper or on a chalkboard, but a computer can be a powerful ally, too. Computers were originally created for numerical calculations, but they can also do symbolic mathematics, including algebra and calculus. In Python, you must import a special library to do symbolic calculations. You can also take advantage of free online resources. In this section, we describe both approaches, then apply symbolic computing to our first passage problem.

10.3.1 Wolfram Alpha

For some problems in symbolic computing, a free online resource like wolframalpha.com is the simplest solution. Wolfram Alpha is particularly convenient for “one line” calculations to obtain a formula: evaluating an integral, summing a series, taking a derivative, getting the roots of a cubic or quartic equation, and so on. Go to Wolfram Alpha, and try the examples below. Wolfram Alpha doesn’t use the same syntax as Python, but its ability to interpret natural language is pretty good, so you don’t have to learn a new programming language.

Integrals

Here is an integral that arises in probability theory:

$$\int_{-\infty}^{\infty} dx \frac{1/\pi}{x^2 + 1} \frac{1/\pi}{(a - x)^2 + 1}.$$

Because the value of this integral depends on the value of a , it defines a function of a . Suppose that you were writing a program to and had to evaluate this integral. You could use the methods of Section 6.7 to evaluate the integral for any particular value of a , but if you have to evaluate the integral for many values of a to make a plot, it could take a while. Maybe there is a simple formula you could use instead.

The integral looks daunting, but try entering it into Wolfram Alpha as

```
integrate 1/pi/(1+x^2) * 1/pi/(1+(x-a)^2) from -oo to oo assuming a>0
```

(Wolfram Alpha recognizes “oo” as a synonym for ∞ .) Notice that Wolfram Alpha was able to interpret the query even though we did not use any special commands. (If you leave off the “assuming $a > 0$ ”, you may not get as useful a result, though.)

Wolfram Alpha returns a simple result. This integral is the *convolution* of two probability density functions. In this case, they are each called Cauchy distributions. Perhaps surprisingly, the result is also of the same general form as the distributions we started with.¹¹

Here is another example. The integral $\int_p^q dx x^n (1 - x)^{M-n}$ arises in the study of **credible intervals** for the parameter x that defines a Bernoulli trial, given that M trials yielded n successes.¹² We can make progress evaluating it by entering

```
integrate x^n * (1-x)^(M-n)
```

This time, we did not specify the range of integration. (You can try to use the general limits p and q , but Wolfram Alpha may not be able to evaluate the definite integral.) Because we don’t specify the range of integration, Wolfram Alpha responds with the indefinite integral, a function of x , M , and n . It may not seem helpful to be told that this function is the “incomplete beta function,” but now we can see if Python knows about that function (even if we don’t). It does: It is the function **betainc** in the **scipy.special** module. We can now write code that evaluates this function instead of computing an integral numerically:

```
from scipy.special import betainc

def credible_interval(p, q, M, n):
    return betainc(n+1, M-n+1, p) - betainc(n+1, M-n+1, q)
```

¹¹ Nelson, 2015, Chapter 5 discusses Cauchy distributions.

¹² Nelson, 2015, Chapter 6 discusses credible intervals, and this example in particular.

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Now you can quickly calculate the result without doing any numerical integration.

If you only need the definite integral from 0 to 1, Wolfram Alpha can evaluate this:

```
integrate x^n * (1-x)^(M-n) from 0 to 1 assuming M>n
```

We learn that this integral simplifies to

$$\frac{\Gamma(n+1)\Gamma(M-n+1)}{\Gamma(M+2)}.$$

This expression is easy to include in Python code: There is a `gamma(x)` function in `scipy.special`.¹³

In general, evaluating a special function is faster and more accurate than numerical integration. However, this option is not always available. If you cannot locate a predefined function corresponding to the expression returned by Wolfram Alpha—or, if Wolfram Alpha does not return anything helpful—you may be able to use `quad` to get the numerical result you need. (See Section 6.7.)

Sums

You may have forgotten the result of the infinite discrete sum $\sum_{k=0}^M k^3$, but typing

```
sum k^3 from 0 to M
```

into Wolfram Alpha tells you it's $M^2(M+1)^2/4$, a result you may want someday for computing the third moment of a uniform discrete probability distribution. If you forget the formula for the geometric series, you can get it just as easily:

```
sum r^k over k from 0 to M
```

This time, we avoided ambiguity by specifying what variable is to be summed (not `r`).

Ordinary differential equations

Wolfram Alpha can find exact solutions to simple differential equations, like the kinematic equation for constant acceleration:

```
solve d^2 x / dt^2 = a
```

Wolfram Alpha can often solve more complex differential equations, too. The equation $dv/dt = -Av + Be^{-ct}$, where A , B , and c are constants, arises in the context of modeling virus dynamics. (See Section 5.1.1.) We can ask Wolfram Alpha to solve it:

```
solve dv/dt = -A v + B e^{(-C t)} with v(0) = D
```

To get a definite solution, we need to specify an initial value. In the expression above, we required that $v(t)$ should equal the constant D at time zero. This solution is where Equation 5.1 came from.

Another ordinary differential equation arising in bacterial dynamics can be solved similarly:¹⁴

```
solve dx/dt = t - x, x(0) = 0
```

This shows where the formula for $W(t)$ in Equation 5.2 came from.

¹³ For integer values of M and n , the gamma function is equal to the more familiar factorial: $\Gamma(n+1) = n!$. The `factorial` function is also available in `scipy.special`.

¹⁴ See Section 5.2.1 (page 73). Nelson, 2015 discusses this system and its physical model.

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10.3.2 The SymPy library

If you need a quick formula, Wolfram Alpha is quite useful. However, it is difficult to incorporate Wolfram Alpha into a Python program. The SymPy library enables Python to do symbolic mathematics.¹⁵ SymPy can also do “infinite precision” numerical calculations and turn symbolic expressions into functions that act on NumPy arrays.

To get some idea of what is possible with SymPy, start by typing the following commands in the IPython console:

```
from sympy import *
init_session()
```

SymPy is a library where `import *` is so convenient that we make an exception to our general advice and bring all its functions into our working environment. When Python executes `init_session()`, it defines several variables and functions. The following examples will not work if you omit this command.

Algebra

SymPy can do symbolic algebra. It uses standard Python notation for arithmetic operations.

```
expand( (x + y)**5 )
factor( x**6 - 1 )
```

If you wish to see the formula for the roots of a cubic, or use the formula in a calculation, SymPy can help:

```
a, b, c, d = symbols('a b c d')
r1, r2, r3 = solve(a * x**3 + b * x**2 + c * x + d, x)
display(r1)
print(r1)
5 latex(r1)
```

In the first line, the function call to `symbols` asks SymPy to generate four new symbols. The right-hand side binds those symbols to Python variable names `a, ..., d`. The string argument tells SymPy how we would like those symbols to be displayed. In this case, we will see italic letters matching the variable names. However, we could instead have asked for Greek names via `symbols('alpha beta gamma delta')`.

Our four new symbols are abstract algebraic quantities, not floating-point numbers, integers, or any other kind of numerical value. You can also create a collection of symbols with subscripts: `x = symbols('x:10')` will bind the variable `x` to a tuple of 10 symbols: $(x_0, x_1, x_2, \dots, x_9)$. You can then access the symbols in the tuple with an index: `x[8]` refers to x_8 . This can be a convenient way to create and refer to a lot of variables when you need them.

Notice how `display(r1)` displays a symbolic expression that looks like the math in textbooks. `print(r1)` displays a plain-text expression—one that could be copied and pasted into a Python script if desired. `latex(r1)` returns a Python string of L^AT_EX commands. `print(latex(r1))` gives a string that can be copied and pasted directly into a L^AT_EX file if desired—or written directly to a file from within a Python script.

Arbitrary precision

SymPy can also carry out numerical calculations to arbitrary precision. Suppose that we want to evaluate the root `r1` for the particular cubic equation $2x^3 - 3x^2 + 4x = 1$. The following commands will substitute

¹⁵ SymPy is part of the Anaconda distribution and can be installed with `conda`. See Appendix A for details.

the coefficients into the general formula, display the exact solution in terms of radicals, and evaluate it numerically to 50 decimal places:

```
myroot = r1.subs( [(a,2), (b,-3), (c,4), (d,-1)] )
display(myroot)
print( myroot.n(50) )
```

Appending `.n()` to a SymPy expression or object will attempt to evaluate the result to 15 significant figures. You can request more or fewer digits by supplying an integer, as in this example.

Lambdify

A symbolic expression can be converted to a Python function by using a function called `lambdify`. For example, the following code will create a function that returns the roots of any cubic equation.

```
solve_cubic = lambdify( [a, b, c, d], [r1, r2, r3] )
```

`lambdify` requires two arguments: a list of symbols that will be the arguments of the new function and a list of SymPy symbols or expressions to evaluate. Either list can be replaced by a single symbol or expression. The resulting function (`solve_cubic` in this example) is now a regular Python function that can be used outside of SymPy. It can even perform vectorized math on NumPy arrays.

```
# Solve 2 * x**3 - 3 * x**2 + 4 * x - 1 = 0.
solve_cubic(2, -3, 4, -1)

# Get roots of 100 random cubic equations of similar form.
5 N = 100
A = np.random.random(N) - 0.5
B = np.random.random(N) - 0.5
C = np.random.random(N) - 0.5
D = np.random.random(N) - 0.5
10 R = solve_cubic(A, B, C, D)
```

This is one of the great benefits of working with SymPy. You can start with a symbolic problem, obtain a symbolic solution, and transform it into a numerical function, all within Python. You don't have to re-type any long formulas that may be generated along the way, which reduces the chance of errors.

Calculus

SymPy can also do limits, derivatives, integrals, and sums:

```
limit( x*log(x), x, 0 )          # Limit of indeterminate form
diff( x*exp(-x**2), x, x, x )    # Third derivative of a function
integrate( cos(x)**2, x )         # Indefinite integral
integrate( exp(-x**2), (x, -oo, oo) ) # Definite integral
5 Sum( k**3, (k, 0, m) ).doit().factor() # Sum a series
```

The case of a function matters in SymPy. Lower-case functions, like the first four commands in this example, will attempt to evaluate the expression. Upper-case functions, like the last command, return an expression without attempting to evaluate or simplify it in any way. The method called `doit()` will attempt to evaluate an expression that involves sums, limits, integrals, and so on. Methods like `factor()`, `expand()`, `collect()`, and `simplify()` can produce equivalent expressions that are

easier to interpret.

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Power series

One useful application of SymPy in numerical work is to obtain a power series approximation to a function that might suffer from numerical error if evaluated naively. This can happen with indeterminate forms like $\sin(x)/x$, but it can also happen with perfectly well-behaved functions. Because floating point numbers are only stored with a finite number of digits (about 16), taking the difference of two very large numbers or two numbers that are nearly equal can lead to large rounding errors. NumPy does not account for this, and numerical error of this kind can spoil a calculation.

As an example, let's use special relativity to calculate the difference between the time that elapses on your wristwatch and the time that elapses on a stationary clock while you go for a stroll.

```
velocity = np.linspace(-3, 3, 101) # walking velocities, in m/s
c = 299792458 # speed of light, in m/s
tdf = 1/np.sqrt(1-v**2/c**2) # time dilation factor
t0 = 1000 # time of walk, in seconds, on your watch
5 t = t0 * tdf # elapsed time on a stationary clock
dt = t - t0 # difference in times
plt.plot(velocity, dt)
```

The time difference only takes on two discrete values, and most of the elements of `dt` are zero! The difference should be small, but not zero. In fact, it is so small that floating point arithmetic can't tell the difference from zero. However, we can use SymPy to create a function to calculate the difference in time for arbitrarily small speeds. We let $x = v/c$ and expand $\frac{1}{\sqrt{1-x^2}} - 1$ in a Taylor series around $x = 0$:

```
f = series(1/sqrt(1-x**2) - 1, x=0, n=7)
display(f)
delta = lambify(x, f.remove0())
```

Now we can recalculate the difference in times and plot.

```
dt = t0 * delta(velocity/c)
plt.plot(velocity, dt)
```

(`f.remove0()` in Line 3 above tells SymPy to discard the $\mathcal{O}(x^7)$ piece of the expansion before turning it into a function for numbers and arrays.) We see that the difference is small—on the order of a few femtoseconds—but not zero. We also see that the difference is a smooth function of the velocity, and that the direct numerical approach did not get a single value correct!

Keep in mind, however, that a series expansion is only valid when its argument is small.

Differential equations

SymPy can also solve differential equations.

```
result = dsolve( diff(f(x),x) + f(x) - x, f(x) )
display(result)
display(result.rhs)
```

A call to `dsolve` returns an object called an *Equality*. A SymPy expression for the solution is contained in the right-hand side of this equality and can be accessed as shown in this example.

You can specify initial conditions (or other boundary values) by passing a Python dictionary with the `ics` keyword argument. This allows you to use the solution with other Python code. The following

example illustrates how to solve a differential equation with SymPy and plot the result with PyPlot:

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```

# Use SymPy to solve a differential equation.
A = symbols('A')
result = dsolve( diff(f(x),x) + f(x) - x, f(x), ics={f(0):A})
F = lambdify([A, x], result.rhs)

# Plot solutions for different initial conditions.
time = np.linspace(0,10,101)
initial_conditions = np.arange(5)
for f0 in initial_conditions:
    plt.plot( time, F(f0, time), label=r"$f(0) = {:.2f}{}".format(f0) )
plt.legend()

```

Plotting functions

SymPy also has its own plotting library. It is useful for making quick plots from the command line:

```
plot( besselj(0, x), besselj(1, x), sin(x)/x, (x, 0, 10) )
```

SymPy's plotting commands are rather different from PyPlot, and we will not discuss them in detail. Use `help(plot)` to learn more.

10.3.3 Other alternatives

SymPy is useful for many quick symbolic calculations and plots. At present, however, its ability to evaluate integrals and solve differential equations is not as good as Wolfram Alpha's. If you are doing extensive symbolic computation, Wolfram Alpha may not be adequate, either. Commercial software like *Mathematica* or Maple, or the free alternative Sage may be better choices.¹⁶ Despite its limitations, SymPy is useful for its ability to bridge the worlds of symbolic mathematics and numerical computation within the Python ecosystem.

For more information about SymPy, visit docs.sympy.org/latest. Its documentation is also extensive and filled with examples.

10.3.4 First passage revisited

Let's now apply some of these computer algebra tools to the first passage case study introduced earlier. When you ran the simulations in the first two sections of this chapter, you may have noticed that the histograms of first passage times suggest a power law distribution.¹⁷ The number of random walkers that take k steps to reach distance L is roughly proportional to $k^{-3/2}$. When a pattern like this emerges from a simulation, we should ask if it is a coincidence or if there is a deeper reason for it. These “deeper reasons” often give us insight into the system and allow us to make connections to other models and systems.

Probability distribution

First, we might ask ourselves about the probability distribution of locations for a random walker after

¹⁶ Sage uses Python to bring together over 100 open-source mathematics libraries (www.sagemath.org). It is a large and powerful computer algebra system. While it is possible to import Sage into a Python program, Sage is more commonly used as a standalone interactive program or scripting language.

¹⁷ See Your Turn 10B (page 135).

taking k steps. You explored this problem in Chapter 7, where your data suggested that the mean square displacement after k steps was equal to k :

$$\langle x_k^2 \rangle = k.$$

For an unbiased random walk, we are just as likely to end up an equal distance to the left or the right of our starting position, so the average displacement after k steps should be zero:

$$\mu_k = \langle x_k \rangle = 0.$$

If you think back to a statistics course or an introduction to statistic in a lab course, you might remember two relevant bits of information:

- The expectation and standard deviation are useful statistical quantities.
- A “normal” distribution (sometimes called “Gaussian”) is completely characterized by its expectation and standard deviation.

We have not yet calculated the standard deviation for the location after k steps, but we can:

$$\sigma_k^2 = \langle x_k^2 \rangle - \langle x_k \rangle^2 = \langle x_k^2 \rangle - k = k.$$

We can construct a probability density function for being at location x after k steps by using the mean and standard deviation:

$$p(x, k) = \frac{1}{\sigma_k \sqrt{2\pi}} e^{-(x-\mu_k)^2/(2\sigma_k^2)} = \frac{1}{\sqrt{2k\pi}} e^{-x^2/(2k)}. \quad (10.1)$$

Equation 10.1 is a working hypothesis, not a mathematical derivation. Plotting this function together with histograms of random walks shows that it describes the data pretty well, and a more formal analysis shows that this is the distribution we expect for a random walk. (See Feynman et al., 2010, Chapter 6.)

Equation 10.1 does not explain the trend in the first-passage data, because $p(L, k)$ is proportional to $1/\sqrt{k}$, whereas we observed that the distribution of first passage times scales as $k^{-3/2}$. Nevertheless, we will soon see that this function plays a role in the true solution, so let’s implement it in SymPy:

```
x, k = symbols('x k', positive=True)
p = exp(-x**2 / 2 / k) / sqrt(2*k*pi)
f = lambify([x, k], p)
```

$f(x, k)$ is now a Python function we can use with NumPy arrays to compare this model with our numerical simulations.

If we imagine a random walker taking steps at regular intervals, then the number of steps k is a measure of time. If many walkers are released at $x = 0$, then we can interpret $p(x, k)$ as the fraction of walkers that are at location x after time k . As k increases, the number of walkers at location $x = L$ will rise to a maximum, then fall. The time $k_*(L)$ when that peak occurs might be relevant to when a single walker is most likely to first arrive at $x = L$. Let’s explore this idea by taking the derivative of p with respect to k , setting it equal to 0, and solving for k :

```
dp = diff(p,k).simplify()
display(dp)
solve(dp, k)
```

We see that the peak probability occurs after $k_* = L^2$ steps. Tantalizingly, we also see that the derivative has a piece that scales as $1/k^{3/2}$ and a piece that scales as $x^2/k^{5/2}$. Both terms are multiplied with an exponential factor, $e^{-x^2/(2k)}$. However, for very large values $k \gg x^2$, the exponential term will be approximately 1, and the $1/k^{3/2}$ piece will be much larger than the $x^2/k^{5/2}$ piece. So perhaps the *derivative* of our probability distribution holds the key to explaining the distribution of first passage times.

Our hunch also makes qualitative sense: The histogram of first passage times shows the number that arrived at $x = L$ in some time interval—the number of walkers that reached $x = L$ for k between the edges of that interval. The histogram bins are therefore telling us about the *rate* at which particles arrive, not the probability of being there.

To follow up on this hunch, we can turn to a physical model that is closely related to our mathematical process: diffusion. Diffusion can be thought of as a continuum limit of certain kinds of random walks; conversely, a random walk can be interpreted as a series of snapshots of diffusing particles. In one dimension, the diffusion equation for the concentration $c(x, t)$ of particles at position x after time t is

$$\frac{\partial c}{\partial t} - D \frac{\partial^2 c}{\partial x^2} = 0.$$

In fact, Equation 10.1 is a solution of the diffusion equation if we make the substitution $k = 2Dt$, where D is the diffusion constant and t is the elapsed time.¹⁸ Let's verify that claim:

```
D, t = symbols('D t', positive=True)
c = p.subs(k, 2*D*t).simplify()
display(c)
diff_eq = diff(c, t) - D * diff(c, x, x)
5 display(diff_eq)
display( diff_eq.simplify() )
```

We can now reframe our first passage problem as follows: We release a droplet of ink at $x = 0$ and at time $t = 0$ in a narrow tube. The particles diffuse in time, spreading through the tube. The tube ends at $x = L$, but extends infinitely in the other direction. When particles reach $x = L$ they leave, never to return. We already argued that we need to calculate the *current* at $x = L$: the number of particles passing by per unit time. The current is driven by concentration gradients:¹⁹ $I(x, t) = -D\partial c/\partial x$. We can calculate the current as follows:

```
L = symbols('L', positive=True)
I = -D*diff(c, x).simplify()
display( I.subs(x, L) )
```

The current indeed scales as $1/t^{3/2}$! This model makes other predictions that we can test with numerical simulations, too. For example, the current at $x = L$ is proportional to L when k is much larger than L^2 .

We still don't have a complete derivation, though. We have to satisfy boundary conditions. The diffusion equation accounts for particles moving to the left and right with equal probability. Some of the particles contributing to the current at $x = L$ have already passed $x = L$, wandered back into the tube, and are

¹⁸ The role of units in this statement may be puzzling. The diffusion constant is $D = \Delta x^2/(2\Delta t)$, where Δx^2 is the mean square displacement after a single step, and Δt is the average time between steps; thus, D has SI units m^2/s . We have been assuming steps with $|\Delta x| = 1$, ignoring the units. The proper translation from our random walk discussion to physical diffusion is to replace x with $x/\Delta x$ and k with $t/\Delta t$. The factor of 2 is part of the traditional definition of D .

¹⁹ This relation is sometimes called “Fick’s law.”

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passing through for a second time—or a third time, and so on. We need a solution in which any particle that reaches $x = L$ disappears. That is, we require that $p(x, k) = 0$ at $x = L$.

We can satisfy the boundary condition by using the “method of images.” You may have seen this technique when you studied waves in an introductory physics class. When we have a linear differential equation—like the wave equation, the Poisson equation, the Schrödinger equation, or the diffusion equation—we can add together two solutions to get another valid solution. If we are clever, we can pick the two solutions so that they solve our problem in the region of interest and satisfy all of the relevant boundary conditions.

With waves on a string, for example, the amplitude of the wave must be zero at a hard wall. What happens when a pulse is moving toward a boundary at $x = L$? Mathematically, we can satisfy the boundary condition by combining our pulse with another pulse that is inverted relative to $x = L$ and moving in the opposite direction. That is, we take our pulse $f(ct - x)$, which satisfies the wave equation, and add to it an inverted pulse approaching from beyond the boundary: $-f(ct + (x - 2L))$. The combination, $u(x, t) = f(ct - x) - f(ct + (x - 2L))$, is then a solution of the wave equation and satisfies the boundary condition $u(L, t) = 0$. Mathematical theorems of existence and uniqueness guarantee that, not only is this a solution of the wave equation; it is the *only* solution of the wave equation that satisfies our boundary conditions. (See Feynman et al., 2010, Chapter 49 for more details.)

We can adapt this method to the diffusion equation and apply it to our first passage problem. We could get rid of all of the particles that reach $x = L$ by annihilating them. We imagine a droplet of “anti-ink” released at $x = 2L$ at time $t = 0$. When the ink antiparticles collide with ink particles at $x = L$, they annihilate. Because the ink and anti-ink distributions were mirror images to start with and they obey the same diffusion equation, *all* of the particles reaching $x = L$ annihilate with their counterparts, giving $p = 0$ there at all times. Because p is proportional to the concentration, the complete solution of the diffusion equation that models our first passage problem and the associated current at $x = L$ are

$$q(x, t) = c(x, t) - c(2L - x, t) \quad I(L, t) = -D \frac{\partial q}{\partial x} \Big|_{x=L}.$$

We can explore these functions in SymPy:

```
q = c - c.subs(x, 2*L-x)
display(q.simplify())
I = -D*diff(q,x).subs(x,L).simplify()
display(I)
```

The current is then $-D\partial c/\partial x$, called `I` in the code, or

$$\frac{Le^{-\frac{L^2}{4Dt}}}{2\sqrt{\pi Dt^3}}.$$

(The `LATEX` source that we used to typeset this equation was generated directly via `print(latex(I))`.)

At last we have a complete analytic prediction for first-passage times. To compare with the simulation, export it as a Python function:

```
result = I.subs({D: Rational(1,2), t: k})
display(result)
p_first = lambdify([L,k], result)
```

and graph it alongside the simulation data.

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10.4 WRITING YOUR OWN CLASSES

We hope the preceding foray into the first passage problem has illustrated how mathematical models, numerical simulations, data analysis, analytic derivations, and physical reasoning all work together to provide insight into a problem of interest. Python offers a single environment for combining all of these tools. We will now offer a vista onto the next level of coding as we continue to explore first passage.

A huge number of useful packages and libraries are available for Python. But if you program for long enough, you will eventually find a problem for which no library, module, or function exists that precisely meets your needs. You have already learned how to write your own functions and modules. You can also write your own classes—more complex objects with their own data and methods.

Many programming approaches can solve the same problem: imperative, declarative, procedural, functional, recursive, object-oriented. (You are not expected to know the meanings of these terms. The fact that they exist and mean different things makes the point!) Since most programming languages are mathematically equivalent to an abstract device called a “universal Turing machine,” we cannot say that one approach is intrinsically superior to another. However, one may be better suited to a particular problem—or to your way of thinking about that problem—than another.

Writing your own classes is the centerpiece of **object-oriented programming**. You won’t need this approach for many everyday tasks. But when a project grows past a certain complexity, using classes and object-oriented programming is worth the effort. Many textbook examples and homework assignments do not exceed this level, and so it can be hard to see why you should go to the effort of writing your own classes. We close with an investigation of universal behavior in random walks that illustrates the value of an object-oriented approach.

Universal behavior?

We have investigated the statistical properties of random walks and explored interesting behavior in the first passage problem. We have seen how an analytic model of the random walk can yield insight into these numerical simulations. But we have only looked at two kinds of random walks: steps of equal size to the right or left in one dimension, and steps of equal size on a two-dimensional square lattice. This raises several questions. Are the results we have seen particular to these models? Or are they more general? What happens if the step sizes are not equal? What happens when a random walker explores a different kind of lattice? What happens in three dimensions? Are there mathematical results that carry over into more abstract spaces of N dimensions?

Let’s investigate these questions by creating Python objects to simulate a variety of random walks.

10.4.1 A random walk class

We want to look at many different types of random walks but answer the same basic questions about all of them. We can start with two very general questions: (1) What do random walks from different models *look* like? (2) Do they have different statistical properties? We can create a collection of related objects that all have similar structure, data, and methods—a **class** of related objects—to explore these questions.

Rather than build everything up one line at a time, let’s dissect the following code to see what is happening.²⁰ It creates a `RandomWalk` class that does everything *but* create a random walk! We will soon

²⁰ The `nd_random_walk.py` script in the code samples contains all of the following code fragments with more extensive commenting and documentation.

see why implementing this last feature has been postponed.

```
# nd_random_walks.py      [get code]
import numpy as np

class RandomWalk:
    def __init__(self, dimension=1):
        # Create a random number generator.
        self.rng = np.random.default_rng()
        # Store dimension and starting point.
        self.D = dimension
        self.r0 = np.zeros((self.D, 1))

    def _get_steps(self, N):
        # Internal method. Generate individual steps of the random walk.
        return np.zeros((self.D, N)) # Don't go anywhere.

    def get_walk(self, N):
        return np.append(self.r0, np.cumsum(self._get_steps(N), axis=1), axis=1)

    def get_endpoints(self, M, N):
        return np.array([self.get_walk(N)[:, -1] for m in range(M)]).transpose()

    def get_distances(self, M, N):
        return np.sqrt(np.sum(self.get_endpoints(M, N)**2, axis=0))
```

The preceding code sample introduces several new features:

- 1. class statement:** Constructing a class starts with the `class` statement, followed by the name of the class (`RandomWalk`), followed by a colon. The indented blocks on the following lines are all part of the class definition. These function definitions are *methods* of the `RandomWalk` class.
- 2. self:** You will almost never see the variable name `self` outside of a class definition. Within a class definition, it refers to the object being created. *Every function definition must include self as its first argument.* No other arguments are required, though they are often useful. When you call the methods of an object, you will not provide any argument for `self`. In fact, Python will raise an exception if you do. We will see examples below, but you can interpret this behavior as follows:

An object always passes itself as the first argument to any of its own methods.

Within the function definitions, we see many references to `self`. For example, `self.D` and `self.r0` are *data fields* of the `RandomWalk` class. Each time we create an object of this class, it will store its own dimension (`self.D`) and starting point (`self.r0`). These attributes are similar to the `size` and `shape` attributes of an `ndarray`. We also see function calls that involve `self`, such as `self._get_steps(N)` and `self.get_walk(N)`. These are calling the `get_steps` and `get_walk` methods of the `RandomWalk` object.

When you first start creating classes, it is easy to forget `self` and end up with errors and exceptions.

- 3. __init__ method:** We can define as few or as many methods for a class as we please. However, some come standard with every Python class, and `__init__` is one of these. It is called a **constructor**,

and it is the function that will be called every time you create a new instance of the `RandomWalk` class. By default, this function will do nothing, and if you do not need to do anything special when you create an object, you do not need to define an `__init__` function. Here, we plan to explore random walks in different dimensions, and the shapes of arrays depend on which dimension we are looking at. Thus, we write our own constructor function. It allows the user to provide a dimension and uses the default value $D = 1$ if none is provided. It then stores the dimension and an array for the starting point of a random walk in `self.D` and `self.r0` respectively. The constructor also gives each `RandomWalk` object its own internal random number generator.

- 4. Private and public methods:** You may wonder why `__init__` and `_get_steps` are preceded by underscores and `get_walk`, `get_endpoints`, and `get_distances` are not. This is Python's convention for distinguishing private and public methods. Private methods are intended to be used internally, not called from the command line or used in scripts. Public methods are approved for general use.²¹

This code is just the class definition. It does not create any new objects. After Python has executed this definition, typing `RandomWalk()` will create a new object from the class named `RandomWalk`. The new object is independent of any other objects created earlier or later by `RandomWalk()`. It's called an `instance` of the `RandomWalk` class. If we want to use this object, we should assign it to a variable: `walk=RandomWalk()`. Now we can access all of the data and methods of the `RandomWalk` instance through `walk`.²²

You can already use this new class, even though the results are not very interesting.

```
import nd_random_walks as RW
import matplotlib.pyplot as plt

walk_1d = RW.RandomWalk()
5 walk_2d = RW.RandomWalk(2)
walk_3d = RW.RandomWalk(dimension=3)

fig, ax = plt.subplots(1,2)
ax[0].plot(*walk_1d.get_walk(100))
10 ax[1].hist(walk_2d.get_distances(10, 100), bins=10, range=(0,100))
print("The {}-D random walk starts at\n{}".format(walk_3d.D, walk_3d.r0))
```

The shape of the arrays generated by this class allows use of the `*args` syntax described in Section 10.1.2 for plotting. `plt.plot(*walk.get_walk(N))` will plot the random walk in 1, 2, or 3 dimensions.

Neither the time series nor the histogram is very interesting. This example merely illustrates how to create and interact with objects from our random walk class. As we introduce more examples, we recommend that you build up the random walk classes in a script of your own. You can also run the `nd_random_walks.py` script from this book's code repository.²³ If you are working at an IPython command line, with Jupyter, or with Spyder, remember to `reload` the module every time you make changes, or the changes will not take effect in your session.²⁴

²¹ Some languages, like C++, allow for truly private functions that *cannot* be called in other code. Python has no such inaccessible functions, but the convention described here forces you to recognize what you are doing by typing an awkward underscore or two before the function name.

²² We used this mechanism with linear regression in Section 10.2.2 (page 135).

²³ Code is available starting from press.princeton.edu/titles/32489.html.

²⁴ See Section 6.1.2.

Now let's introduce some more interesting random walks by building on top of this `RandomWalk` class, using **inheritance**. All of our new random walk classes will "inherit" the data and methods of the `RandomWalk` class. We can then add the extra features we want.

Let's start with walks on a "square" lattice in any dimension.

```
5   class LatticeWalk(RandomWalk):
6       def __init__(self, dimension=1):
7           super().__init__(dimension)
8           # Create list of legal steps on D-dimensional square lattice.
9           M = np.eye(self.D) # Get column vectors from identity matrix.
10          self.basis = [ M[:,n] for n in range(self.D) ]
11          self.basis += [ -M[:,n] for n in range(self.D) ]
12
13      def _get_steps(self, N):
14          # Generate individual steps by randomly choosing from self.basis.
15          return self.rng.choice(self.basis, size=N).transpose()
```

For $D = 1$, this will be the original random walk we studied: steps of equal length, to the right or left with equal probability. For $D > 1$, this class will take steps of equal length along one of the $2D$ allowed directions. (In two dimensions, these would be up, down, left, or right.)

Before we start exploring random walks on 11-dimensional hyperlattices, let's look at some important features of this second class definition.

- 1. Inheritance:** The `class` statement is different this time. `LatticeWalk(RandomWalk)` looks like a function call, but in a `class` statement, it tell Python, "A `LatticeWalk` is a particular kind of `RandomWalk`." Thus, a `LatticeWalk` automatically has access to all of the data and methods defined in the `RandomWalk` class. It will also have whatever additional data and methods we define here. What is more, any data fields or methods we define here will override those in the "parent" class `RandomWalk`. This means, for example, that a `LatticeWalk` will have the `get_walk`, `get_endpoints`, and `get_distances` methods defined in the `RandomWalk` class, but it will define its own constructor and `_get_steps` method.
- 2. `super()`:** This function refers to the parent class and allows us to access *its* data and methods, even if the current class has attributes with the same names. Here, `super()` is equivalent to `RandomWalk`. Line 3 could be replaced by `RandomWalk.__init__(dimension)` with no change in behavior. We use `super()` to simplify code maintenance. If we change the name of the `RandomWalk` class, or to make a `LatticeWalk` inherit data and methods from a different parent class, we only have to change one occurrence of "RandomWalk" and `super()` will take care of the rest.
- 3. Descent with modification:** The `LatticeWalk` constructor calls the `RandomWalk` constructor first, then does some additional work. It defines a list of allowed steps and stores it as `self.basis`. The `LatticeWalk` class also overrides the `_get_steps` method of `RandomWalk`. Instead of returning an array of zeros, it uses its random number generator (defined in the `RandomWalk` constructor) to randomly choose N of its allowed steps.

Notice how much of the `RandomWalk` code was reused. We have a consistent interface for many different kinds of random walks. Now let's put it to work!


```

from importlib import reload
reload(RW)      # Use if nd_random_walk.py was modified since last import.

lattice_1d = RW.LatticeWalk(dimension=1)
5 lattice_2d = RW.LatticeWalk(dimension=2)
lattice_3d = RW.LatticeWalk(dimension=3)

N = 50  # number of steps in each walk
M = 5   # number of walks to plot
10
# Plot 3 random walks of each kind.
from mpl_toolkits.mplot3d import Axes3D
fig = plt.figure()

15 ax1 = fig.add_subplot(1, 3, 1)
for m in range(M): ax1.plot( *lattice_1d.get_walk(N) )
ax1.set_title('1-D Time Series')

ax2 = fig.add_subplot(1, 3, 2)
20 for m in range(M): ax2.plot( *lattice_2d.get_walk(N) )
ax2.set_title('2-D Random Walks')

ax3 = fig.add_subplot(1, 3, 3, projection='3d')
25 for m in range(M): ax3.plot( *lattice_3d.get_walk(N) )
ax3.set_title('3-D Random Walks!')

# Look at statistics for many walks.
N = 100      # number of steps in each walk
M = 10**5    # number of walks to analyze
30
fig, ax = plt.subplots(1,3)

ax[0].hist(lattice_1d.get_distances(M,N), bins=20)
ax[0].set_title('1-D Distance Distribution')
35
ax[1].hist(lattice_2d.get_distances(M,N), bins=20)
ax[1].set_title('2-D Distance Distribution')

ax[2].hist(lattice_3d.get_distances(M,N), bins=20)
40 ax[2].set_title('3-D Distance Distribution')

```

Some interesting behavior is emerging. Even though the three walks have the same number of steps and all of the steps are the same length, walkers on a cubic lattice end up farther away than walkers on a square lattice or a line. The shape of the probability distribution changes, too.

What about other lattices? A `LatticeWalk` chooses each step from a list of allowed steps. If we define the allowed steps properly, we can simulate walks on any lattice in any dimension. If we use inheritance, we can do so with very little extra code. Let's add two-dimensional triangular and honeycomb lattices to our growing collection of random walks.

```

class TriangularWalk(LatticeWalk):
    def __init__(self, dimension=2):
        super().__init__(dimension=2)
        # Create list of legal steps on a triangular lattice.
        5      cosTheta, sinTheta = np.cos(np.pi/3), np.sin(np.pi/3)
        R = np.array([[cosTheta, -sinTheta], [sinTheta, cosTheta]])
        v = np.array([0,1])
        self.basis = []
        for n in range(6):
            10     self.basis += [v]
            v = np.dot(R,v)

class HoneycombWalk(LatticeWalk):
    def __init__(self, dimension=2):
        super().__init__(dimension=2)
        # Create list of legal steps on a honeycomb lattice. See _get_steps().
        15     cosTheta, sinTheta = np.cos(2*np.pi/3), np.sin(2*np.pi/3)
        R = np.array([[cosTheta, -sinTheta], [sinTheta, cosTheta]])
        v = np.array([0,1])
        self.basis = []
        20     for n in range(3):
            self.basis += [v]
            v = np.dot(R,v)

25     def _get_steps(self, N):
        # Honeycomb lattice has two sublattices. (-1)**n accounts for this.
        return self.rng.choice(self.basis, size=N).T * (-1)**np.arange(N)

```

The `TriangularWalk` and `HoneycombWalk` classes inherit from the `LatticeWalk` class, which inherits from the `RandomWalk` class. Again, we only have to define what is different in the new classes: a new list in `self.basis` for both, and a modified `_get_steps` method for the honeycomb lattice.²⁵

```

reload(RW) # Use if nd_random_walk.py was modified since last import.

# Store RandomWalk objects in a dictionary whose keys are plot titles.
walkers = { "Square Lattice": RW.LatticeWalk(dimension=2),
            5           "Triangular Lattice": RW.TriangularWalk(),
            "Honeycomb Lattice": RW.HoneycombWalk() }

# Plot samples of each type.
N = 50 # number of steps in each walk
10    M = 5 # number of walks to plot

fig, ax = plt.subplots(1,3)
for n,k in enumerate(walkers.keys()):
    15    for m in range(M): ax[n].plot(*walkers[k].get_walk(N))
    ax[n].set_title(k)

```

²⁵ Technically, the honeycomb “lattice” is a triangular lattice with a basis.

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```
# Look at statistics for many walks.  
N = 100      # number of steps in each walk  
M = 10**5     # number of walks to analyze  
  
20 fig, ax = plt.subplots(1,3)  
for n, k in enumerate(walkers.keys()):  
    ax[n].hist(walkers[k].get_distances(M,N), bins=40)  
    ax[n].set_title(k)
```

We see that the individual walks look rather different, but the distribution of distances looks rather similar.

As a final variation on the random walk theme, we consider walks of variable step size in random directions. We can still use the basic `RandomWalk` class to define the interface, but now we must introduce new methods to generate steps. The following classes will generate random walks in any dimension.

```
class DirectionalWalk(RandomWalk):  
    def _direction(self, N):  
        # Use the Box-Muller transform to generate N random unit vectors.  
        vectors = np.random.normal(size=(self.D,N))  
        lengths = np.sqrt(np.sum(vectors**2,0))  
        return vectors/lengths  
  
    def _magnitude(self, N):  
        # Only the direction is random for this class.  
        return np.ones(size=N)  
  
    def _get_steps(self, N):  
        # Return coordinates for random walk of N steps in D dimensions.  
        return self._direction(N) * self._magnitude(N)  
  
15 class UniformWalk(DirectionalWalk):  
    def _magnitude(self, N):  
        # Draw N step sizes from the uniform distribution. Mean size is 1.  
        return 2*self.rng.random(size=N)  
  
20 class GaussianWalk(DirectionalWalk):  
    def _magnitude(self, N):  
        # Draw N step sizes from chi distribution. Mean size is 1.  
        return np.sqrt(self.rng.chisquare(df=self.D, size=N)/self.D)  
  
25 class ExponentialWalk(DirectionalWalk):  
    def _magnitude(self, N):  
        # Draw N steps from exponential distribution. Mean size is 1.  
        return self.rng.exponential(size=N)  
  
30 class ParetoWalk(DirectionalWalk):  
    def __init__(self, dimension=1, nu=2):  
        super().__init__(dimension)  
        self.nu = nu
```

35 self.norm = max(0.01,nu-1)

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```
def _magnitude(self, N):
    # Draw N step sizes from Pareto distribution. Mean is 1 if nu > 1.01.
    return self.rng.pareto(a=self.nu, size=N) * self.norm
```

The `DirectionalWalk` class generates steps of uniform length in a random direction. The other classes add a variable step size to this class. The common interface makes it easy to compare different walks.

```
reload(RW) # Use if nd_random_walk.py was modified since last import.

# Store RandomWalk objects in a dictionary whose keys are plot titles.
walkers = { "Directional": RW.DirectionalWalk(dimension=2),
            "Uniform": RW.UniformWalk(dimension=2),
            "Gaussian": RW.GaussianWalk(dimension=2),
            "Exponential": RW.ExponentialWalk(dimension=2),
            r"Pareto $\nu=2$": RW.ParetoWalk(dimension=2, nu=2),
            r"Pareto $\nu=4$": RW.ParetoWalk(dimension=2, nu=4) }

# Plot samples of each type.
N = 50 # number of steps in each walk
M = 5 # number of walks to plot

fig, ax = plt.subplots(2,3)
for n,k in enumerate(walkers.keys()):
    I, J = n // 3, n % 3 # Get subplot indices.
    for m in range(M): ax[I,J].plot(*walkers[k].get_walk(N))
    ax[I,J].set_title(k)

# Look at statistics for many walks.
N = 100 # number of steps in each walk
M = 10**5 # number of walks to analyze

fig, ax = plt.subplots(2,3)
for n,k in enumerate(walkers.keys()):
    I, J = n // 3, n % 3 # Get subplot indices.
    ax[I,J].hist(walkers[k].get_distances(M,N), bins=40)
    ax[I,J].set_title(k)
```

If you run this code and create the plots and histograms, you will see that the behavior of the random walks is quite similar—except for the `ParetoWalk` class. We leave it to you to explore this in detail!

10.4.2 When to use classes

This random walk study is well suited to object-oriented programming, although it could be explored in many ways. There are many programming styles, and none is ideal for every problem. Problems with the following properties lend themselves to object-oriented programming:

- **Inheritance.** When the problem can be organized into a hierarchy of objects that have many similar data fields and methods—like the random walks we explored—you can use classes to organize your code. *Define once, and reuse often.*

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- **Encapsulation.** Sometimes different instances of similar objects require their own private data and methods for transforming that data. Rather than keep many separate lists and specialized functions “out in the open,” risking name collisions, we can collect associated data and functions into individual objects—*encapsulating* it. (Imagine working with 12 arrays and trying to keep track of `size1`, `size2`, ..., `size12`, `shape1`, `shape2`, ..., `shape12`, and `data1`, `data2`, ..., `data12`!) Organizing data into dictionaries is sufficient in many cases. Classes provide even more control and flexibility.
- **Separate interface and implementation.** In many cases, people only need to know how to use certain objects, not how they work, in order to accomplish their tasks. (Think of the `ndarray` of NumPy, or the `Figure` and `Axes` objects of PyPlot, or indeed your car or web browser.) Classes and object-oriented programming allow you to define an interface—a way for people to use your objects—without specifying the details of implementation. You can organize the data and write the code inside the functions however you like, and you can make changes that improve performance without affecting the way people *use* your software.
- **Object-oriented problems.** Sometimes we think about problems in terms of objects interacting with one another. Writing code that reflects the way you think about a problem is often easier than changing the way you think about a problem to write code. For example, if you were simulating an optical system, you might think of it in terms of rays that come from sources and get reflected by mirrors and refracted by lenses. You could try to solve the lens and mirror equations for this system and code up the resulting formulas, or you could create `Source`, `Ray`, `Lens`, and `Mirror` objects that obey simple rules and investigate what happens in different arrangements. Many complex systems obey simple rules, and modeling a system in terms of interacting objects is an effective strategy for investigating its behavior.

We hope this short introduction to Python classes will allow you to use object-oriented programming when you think it is the right tool for the job.

Get Going

Programming ... can be a good job, but you could make about the same money and be happier running a fast food joint. You're much better off using code as your secret weapon in another profession.... People who can code in biology, medicine, government, sociology, physics, history, and mathematics are respected and can do amazing things to advance those disciplines.

— Zed Shaw (2017)

We hope you have enjoyed getting to know Python. You have learned a lot—enough, we hope, for you to continue learning on your own.

You do not have to study Python to learn more about it. Just apply what you have learned as you study subjects that interest you. When you encounter an unfamiliar mathematical concept (perhaps a “wave packet”), create a plot to gain some intuition. When you find an interesting model, explore its behavior as you vary its parameters. When you come across a new statistical concept (perhaps “skewness,” “kurtosis,” or the “interquartile range”), try applying it to some real or simulated data to learn what it tells you about a collection of numbers. If you do this, then not only will you better understand what you are studying; you will also improve your proficiency with Python. You may need it when it’s time for your own research!

For fun and inspiration, we leave you with a script you can run. Give it a minute—literally. This is an example of what you can do on your own now.

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```
# surprise.py      [get code]
import numpy as np, matplotlib.pyplot as plt
max_iterations = 32
x_min, x_max = -2.5, 1.5
y_min, y_max = -1.5, 1.5
ds = 0.002
X = np.arange(x_min, x_max + ds, ds)
Y = np.arange(y_min, y_max + ds, ds)
data = np.zeros( (X.size, Y.size), dtype='uint')
10 for i in range(X.size):
    for j in range(Y.size):
        x0, y0 = X[i], Y[j]
        x, y = x0, y0
        count = 0
15    while count < max_iterations:
        x, y = (x0 + x*x - y*y, y0 + 2*x*y)
        if (x*x + y*y) > 4.0: break
        count += 1
        data[i, j] = max_iterations - count
20 plt.imshow(data.transpose(), interpolation='nearest', cmap='jet')
plt.axis('off')
```

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APPENDIX A

Installing Python

This appendix explains how to install the Python environment described in the main text. The free Anaconda distribution is provided and maintained by Anaconda, Inc. It includes the Python language, many libraries, the Spyder integrated development environment (IDE), and the Jupyter Notebook application in a single package. (A software **package** is a collection of programs and data files related to a particular application or library. The data files include information about any other packages that may be required for the new software to function properly.) Anaconda also provides a simple way to update packages and install new ones. There are alternatives, however. You may find one better suited to your own needs or preferences. One place to look is python.org.

Our intention is not to promote a particular distribution of Python, but to promote Python itself as a tool for scientific computing.

A.1 INSTALL PYTHON AND SPYDER

You can download Anaconda at anaconda.com/download (“Install Anaconda Individual Edition”).

You must first choose your installer. The site automatically detects your operating system, but do not download anything just yet. First, make sure the installer offered to you matches your operating system: macOS, Windows, or Linux. Second, make sure your operating system meets the requirements for Anaconda. You can find the current requirements at docs.anaconda.com/anaconda/install. Third, make sure you choose the proper version of Python. The instructions below describe how to install the current version of Python 3 at the time of this writing. (Newer versions of Anaconda and Python will presumably follow a similar installation process, but consult the online documentation at docs.anaconda.com if you have trouble.) Finally, you must decide which type of installation program you want: graphical or command line. The graphical installation (point and click) is easiest, unless you already enjoy working with your operating system’s command line.¹

If your computer meets the system requirements and you wish to carry out a graphical installation, proceed to Section A.1.1.

If your operating system does *not* meet these requirements, the standard installation may not succeed on your computer, or the installation may succeed, but some packages may not work properly. You might wish to try Section A.1.1 anyway. If the installation doesn’t work, it is easy to undo the process by uninstalling Anaconda. Another option is to try installing from the command line as described in Section A.1.2. If that also fails, you can download an older version of the Anaconda or Miniconda installer:

Anaconda repo.continuum.io/archive
Miniconda repo.continuum.io/miniconda

Try matching the date of your operating system with the release date of the installer for best results.

¹Appendix B gives a brisk tour of the command line.

The Anaconda website offers extensive documentation, so even if the instructions given here are insufficient, you should still be able to get Anaconda up and running on your computer. For the most detailed and up-to-date installation instructions, go to docs.anaconda.com/anaconda/install/.

A.1.1 Graphical installation

macOS

To get the latest version of Python, find the Download button on anaconda.com/products/individual. Click on the link and then select the “64-Bit Graphical Installer” option under “MacOS.” This is a large download that will install many packages not used in this tutorial; however, the installation is very easy. Unless you are comfortable working from the command line, this is the installation we recommend.

Once the download finishes, find the package and open it. It is probably in your Downloads folder. Double-click on the file to initiate an installation dialog. Unless you want to customize the installation, just click “Continue” until you reach “Installation type.” Click “Change Install Location,” then click “Install for Me Only.” This will create a folder called anaconda3 or Anaconda3 in the opt directory within your home directory and store all the associated files there. Click “Continue.” Finally, click “Install” to initiate a standard installation. You may be prompted for a password.²

Once the installation is complete, you may find a new icon on your Desktop called Anaconda Navigator.app. Open this application. Click on the [LAUNCH] button below “spyder” to start Spyder. You can also start Spyder without Anaconda Navigator.app by double-clicking on spyder in the folder where Anaconda installed it. By default, this is opt/anaconda3/bin or opt/AAnaconda3/bin. A third option is to start Spyder from your operating system’s command line by typing

`spyder`

and hitting <Return>.

Once you have successfully installed Anaconda and launched Spyder, proceed to Section A.2.

Windows

To get the latest stable version of Python, on anaconda.com/products/individual find the “Download” button. Click on the link and then select the “64-Bit Graphical Installer” option under “Windows.” This is a large download that will install many packages not used in this tutorial; however, the installation is very easy. Unless you are comfortable working from the command line, this is the installation we recommend.

Once the download finishes, find the package and open it. It is probably in your Downloads folder. Double-click on the file to initiate an installation dialog. Unless you want to customize the installation, just click “Continue” until you reach “Destination Select.” Select “Install for Me Only” unless you have a good reason for doing otherwise. This will create a folder called Anaconda3 in your home directory and store all the associated files there. Click “Continue” to bring up the “Installation Type” dialog. Click “Install” to initiate a standard installation. You may be prompted for a password.

Once the installation is complete, you may find a new icon on your Desktop called Anaconda Navigator.app. Open this application. Click on the [LAUNCH] button below “spyder” to start Spyder. You can also start Spyder without the Desktop application. After the install, you will find a new folder called Anaconda or Anaconda (64-bit) under Start>All Programs. This folder contains a

² If you ever need to completely uninstall Anaconda from MacOS, just delete the anaconda3 folder and all of its contents.

shortcut to launch Spyder. The script that launches Spyder may also be found in the User directory under `Anaconda3\Scripts\spyder-script.py`; alternatively, `Anaconda3\Scripts\spyder.exe` will also work.

Once you have successfully installed Anaconda and launched Spyder, proceed to Section A.2.

A.1.2 Command line installation

To install Python on a Linux machine, or to install a leaner environment that includes only the packages you want under macOS or Windows, you can use the `conda` package manager. You can directly access this tool from the command line.

In this appendix, “command line” refers to a command line interface for your operating system—not the IPython command prompt. For macOS users, this is `Terminal.app`. For Windows users, it is `cmd.exe` or PowerShell. For Linux users, it is a terminal or bash shell.

To install Anaconda from the command line, you must first download Miniconda. Select the current version of Python 3 for your operating system at docs.conda.io/en/latest/miniconda.html. Once the download is complete, macOS and Linux users will need to run a script. At the command line, type

```
bash ~/Downloads/Miniconda3-latest-MacOSX-x86_64.sh
```

for macOS, or the equivalent command for your operating system, file name, and file location. This will run the script. Windows users should double click the `.exe` file and follow the instructions on the screen. After you agree to the license terms and select a directory for the installation (or agree to the default location), the script will install Python 3 and the `conda` package manager. (Note that you need to specify the complete path: `/Users/username/opt/miniconda3` or `/opt/miniconda3` for macOS or Linux, or `C:\Users\username\Miniconda3` for Windows. If you only provide the name “anaconda,” then files will be installed in a new directory named `anaconda` in your current directory, wherever that may be.)

If this was successful, you can now run Python, but you will not yet be able to use modules like NumPy, PyPlot, or SciPy, and you will not have access to the Spyder IDE. You can now install them, and the entire Anaconda Python distribution, by typing

```
conda update conda
conda install anaconda
```

If you prefer to install individual packages to minimize disk space, this is simple using `conda`: Instead of the second line above, use commands like

```
conda install numpy matplotlib scipy ipython sympy
```

If you watch the screen after you execute these commands, you will see what `conda` does. It determines which packages are necessary to install the specific ones you asked for, downloads all of the required packages, and then installs and links everything.

Python comes with the IDLE integrated development environment. However, if you want to use the Spyder IDE described in this tutorial, you will need to install it.

```
conda install spyder
```

In principle, this should install all the packages you need to run Spyder. Launch Spyder by typing

```
spyder
```


In practice, you may find that some modules are missing and Spyder will not run. Spyder is actually a colossal Python program. When something is amiss, it displays Python error messages. Perhaps Spyder crashes when you try to run it. At the end of the terminal output, you see

```
ImportError: No module named 'docutils'
```

Just issue the command

```
conda install docutils
```

and this problem is fixed. To get Spyder up and running, install any other missing modules using the `conda install` command.

Even after Spyder is running, you may not have access to all of the debugging tools described in this tutorial. If you are writing your own code later and find another missing module, you may be able to install it with `conda`.

To run all of the code samples in this tutorial, you will need to use `conda` to install the following packages:

<code>conda install ipython</code>	IPython interpreter
<code>conda install numpy</code>	NumPy
<code>conda install matplotlib</code>	Matplotlib and PyPlot
<code>conda install scipy</code>	SciPy
<code>conda install pillow</code>	Image processing library
<code>conda install sympy</code>	Symbolic computation library
<code>conda install pandas</code>	Data analysis library
<code>conda install scikit-learn</code>	Machine learning library

To use the Spyder IDE or Jupyter notebooks, you will need to install them separately:

<code>conda install spyder</code>	Spyder IDE (discussed above)
<code>conda install jupyter</code>	Jupyter Notebook

If you install Anaconda, you can run `anaconda-navigator` and `spyder` from the command line. If you use Miniconda instead, you can run these programs from the command line.

This set of packages should suffice to complete the exercises in this tutorial. However, there is a lot more you can do with `conda`. For example, you can set up environments that allow you to switch between Python 2 and Python 3, or use different versions of NumPy and SciPy. You can learn more about `conda` at docs.conda.io/en/latest.

A.2 SETTING UP SPYDER

Now that you have installed Spyder, there are a few settings to fine tune before you start writing and executing scripts. All of these finishing touches will be made using the Preferences. To access the Preferences, click the wrench icon, or select it from a drop-down menu: **Tools>Preferences** on Windows, or **python>Preferences** on macOS. After making the changes below, you may need to restart Spyder before they take effect.

A.2.1 Working directory

You will need to keep track of your files. The easiest way to do this from within Spyder is to tell it to save all of your work in a folder that you specify. Choose “Current working directory” from the list of options

on the left of the Preferences panel. On the right, choose the button next to “the following directory,” and then click on the folder icon to select a directory, or enter the path of the directory you wish to use. You may wish to create a new folder named `scratch` or `current` to work in. After you have selected a folder, click the appropriate buttons so that “Open file” and “New file” use this directory.

You can still access files anywhere on your computer, but setting these options is the easiest way to locate the files you create with Spyder.

A.2.2 Interactive graphics

Spyder’s default option is to display graphics in a “Plots” pane. You can also adjust your Preferences to display plots in the IPython console, similar to programs like Maple and *Mathematica*. However, these plots are static images. You cannot zoom, move, or rotate them. To make interactive plots the default, click the menu item `python>Preferences` and select the “IPython console” tab on the left of the Preferences window. Click on the `[GRAPHICS]` tab at the top of the menu. In the panel called `Graphics backend`, set `Backend` to `Tkinter` or something other than `Inline`. Finish by clicking `[APPLY]` at lower right.

You may have to restart Spyder before this change takes effect.

A.2.3 Script template

It is good practice to include the author, creation date, and a brief description in any scripts you write. You will also import the NumPy and PyPlot modules in nearly every script you write for this tutorial. You can automate most of this by creating a template for files you create with Spyder.

Choose the menu item `python>Preferences` and select the “Editor” tab on the left of the Preferences window. Next, click on the `[ADVANCED SETTINGS]` button at the top of the menu on the right. Click on the button marked `[EDIT TEMPLATE FOR NEW FILES]`. This will open a file in the Editor. Make changes to this file so that it contains text you want to include in every script you write. Here is a sample:

```
# -*- coding: utf-8 -*-
# file_name.py
# Python 3.8
"""

5 Author:      Your Name
Created:      %(date)s
Modified:     %(date)s

Description
10 -----
"""

import numpy as np
import matplotlib.pyplot as plt
```

The expressions with percent signs will insert the date when a file is created. The first line in this sample template is optional. You should include it if your code makes use of any characters not in the “standard ASCII set,” such as non-English characters and accents.

After you have finished editing this file, save your changes and close it.

A.2.4 Restart

After you make these changes and adjust the other preferences to your liking, quit Spyder and relaunch. You should now be ready to carry out any of the exercises in this tutorial.

A.3 KEEPING UP TO DATE

In the future, there will be new releases of Python and its modules, and the people in charge of the Anaconda distribution will work to eliminate bugs. You can get the latest and greatest Anaconda distribution by using the `conda` package manager. This can be done from your operating system's command line.

At the command line, type

```
conda update conda  
conda update --all
```

The package manager will determine which files need to be downloaded, installed, and updated. You can choose to accept or reject the proposed changes, and `conda` will do the rest.

You can update or install individual packages with `conda`. For example, `conda update numpy` will download and install the latest version of NumPy.

The Anaconda Navigator also allows you to update or install individual packages. It provides a graphical interface to the `conda` package manager. Click on the `Environments` tab on the left side of the navigator window. There are drop-down menus and buttons that allow you to manage the packages in your installation. To update a package you have already installed, select `Installed` from the first drop-down menu; to install a new package, select `Not installed` from the first drop-down menu. Locate the package you wish to update or install and click the `Update index ...` button. Then `conda` will take care of the rest.

For more information on `conda`, see docs.conda.io/en/latest.

A.4 INSTALLING FFMPEG

FFmpeg is an open-source encoder for creating audio and video files. It can be used to create animations within Python, as described in Section 8.3. FFmpeg is not part of Python, nor is it part of the Anaconda distribution. However, it can be installed with `conda` just like other Python packages:

```
conda install ffmpeg
```

A.5 INSTALLING IMAGEMAGICK

ImageMagick is a collection of open-source command line tools for creating and manipulating images. It can also be used to create animations within Python, as described in Section 8.3. Like FFmpeg, ImageMagick is not part of the Anaconda distribution, but it can be installed using `conda`.

Many individuals and organizations have made large repositories of code available through the `conda` package manager. A repository like this is called a “channel.” We need to direct `conda` to the appropriate

channel to install ImageMagick. (This appendix illustrates how to access the `conda-forge` channel: conda-forge.org.) Using a channel requires a slight modification to the installation command:

```
conda install --channel conda-forge imagemagick
```

The installation should proceed as with other Python packages, even though ImageMagick is not a Python package. Once `conda` has finished, you can test the installation by asking your operating system to locate the program `magick`. Type

```
which magick
```

at the command line. A message like “`/Users/username/opt/anaconda3/bin/magick`” indicates that Python will now be able to find and run ImageMagick.

You can also download and install ImageMagick by following the instructions at imagemagick.org.

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APPENDIX B

Command Line Tools

This appendix will introduce you to working at the command line. None of this material is essential for programming in Python, but it can make life easier. We focus on three powerful tools:

- the command line
- a text editor
- version control (Git).

This appendix assumes you have installed the Anaconda Python distribution as described in Appendix A.

B.1 THE COMMAND LINE

A command line is a way to access your computer’s operating system without windows, icons, or a mouse—a strictly text-based way to interact with your computer. You can accomplish many common tasks, like finding, creating, and moving files around, with just a few keystrokes.

If you have already worked through some of the exercises in this book, you have probably entered commands at the IPython command prompt. This is similar to a command line, but not the same. A command line gives you direct access to your computer’s operating system; IPython is just one program among many running on your computer.

To access the command line on macOS, launch `Terminal.app`. If you use Linux, launch a terminal or shell. On Windows, launch the Anaconda Prompt application, which is included in the Anaconda distribution. (Windows has other command line options, like `Cmd` and `PowerShell`, but some of the commands in this appendix will not work with these programs.) You should see a mostly empty screen with a solid or blinking cursor next to a prompt. This is the command line. You type commands here, press `<Return>`, and things happen!

Your computer’s file system is a nested hierarchy of folders or directories, each of which can contain both files and other folders. (In this appendix, we use “folders” and “directories” interchangeably.) To see where you are, type `pwd` and press `<Return>`. You should see something like `"/Users/username/"`. This is your **present working directory**—the folder on your computer where your commands will take effect. What files are here? Type `"ls"` and press `<Return>`. This will list the contents of the present working directory.¹ You can look into other directories without changing the default: `ls /Users` will show you the contents of that directory, which include the one you are (still) in.

Other operating system commands allow you run programs, create and edit files, create folders, move files around, and much more. The table below includes a small set of commands that allow you to accomplish a lot.

¹ `ls` suppresses “hidden files,” those whose filename begins with a period. Use `ls -a` to see all the files.

A word of caution: *There is no Undo button at the command line.* Removing or overwriting files and folders will permanently erase them, usually without asking for confirmation.² You cannot find deleted files in the Trash or the Recycle Bin. The FBI may be able to recover these files from your hard drive, but you probably cannot. The commands below are like sharp knives: They are very useful, but you can hurt yourself if you don't use them properly. Using "version control" software like Git gives you a way to undo the effects.³ Frequently backing up your files on an external hard drive or in the cloud is a good idea, too; some of these systems let you "go back in time" to previous versions of a file.

Command	Function	Purpose
<code>pwd</code>	report the present working directory	Where am I right now?
<code>ls</code>	list the contents of the current directory	What's here?
<code>cd</code>	change directory	Go somewhere else.
<code>cp</code>	copy a file	Duplicate a file.
<code>mv</code>	move a file	Transfer a file to another directory or change its name.
<code>rm</code>	remove a file	Destroy a file.
<code>mkdir</code>	make a directory	Create a new folder.
<code>rmdir</code>	remove a directory	Delete an empty folder.
<code>ls -l</code>	list, long	Detailed file list.
<code>cp -r</code>	copy, recursively	Duplicate a directory and all of its contents.
<code>rm -r</code>	remove, recursively	Destroy a directory and all of its contents.

Rather than describe these commands abstractly, let's look at how we can use them to accomplish some useful tasks.

B.1.1 Navigating your file system

Suppose that you download a folder that contains data files you want to analyze using IPython or Jupyter. The easiest way to do this is to start one of these programs from *inside* the data folder. This is difficult with point-and-click methods or the Anaconda Navigator. It is simple from the command line, once you know how to navigate your file system.

The first three commands (`pwd`, `ls`, and `cd`) in the table above are used to navigate and explore your computer's file system. We've already seen how to use `pwd` and `ls` to find out where you are and what files are present there. `cd` allows you to move around (change your present working directory).

When you launch a terminal or the Anaconda Prompt application, it will always start in your "home" directory. Type `ls` to see what is here. You will see a list of files and directories. Your terminal may use color to indicate which is which. File names usually end with an "extension" like `.txt` or `.py`; directory names usually do not have extensions.

Your Downloads folder is usually in your home directory. Move into it and look around: Type `cd Downloads` to move into the directory. Type `pwd` to confirm that your location has changed, and `ls` to see what is here. If there are other directories here—perhaps the collection of data files or code samples for this book—you can move into and investigate them the same way. Try it. You can't break

² You can force commands to ask for confirmation before deleting or overwriting files until you are comfortable working at the command line. Just use `cp -i`, `mv -i`, and `rm -i` in place of the commands given.

³ Section B.3 introduces Git.

anything with these commands.

As you move around, you start to see the structure of your computer's file system. One directory contains files and other directories, which may themselves contain files and other directories, and so on. You can navigate to any file on your computer by typing the chain of directories that contain it, and ending with the file name. This is called the **path** of the file: for example,⁴

```
/Users/username/Downloads/temp/some_script.py
```

You can use the path to locate a file from any other directory on your computer, because it contains instructions starting from the **root directory**, whose name is simply “/”. This is the directory on your computer that contains every file (and every other directory) on your computer. It is not the same as your home directory, and it contains a lot of folders that should be managed by your operating system. You can look around here, but don't make any changes. Your home directory contains most of the files you will directly create and access.

You can always get back to your home directory easily: Just type `cd` with no arguments. (Then use `pwd` to confirm your location.) You can also type `cd ~` because “~” is a nickname for the home directory.

As you are navigating your file system, you might not want to go all the way back to the home directory. You can use `cd ..` to move up one level in the hierarchy of files, from your current directory into the directory that contains it. That is, “..” is a special symbol whose meaning changes as you move around. It always refers the directory containing the one you are in. Try typing `cd ..` and `pwd` a few times to see how this works. You may end up in your root directory, but you know how to get back home.

You can use `ls` to look in other directories without moving into them, or to look for specific types of files. Move to your home directory and type `ls Downloads`. You will see the contents of the Downloads folder, even though you are in your home directory. You can instead look for files whose names match patterns. For example, `ls Downloads/*.py` will list all Python scripts in your Downloads folder. Similarly, `ls Downloads/*.pdf` will list all of the PDF files, and nothing else. It works because “*” is a **wildcard** character used to match patterns. Thus, `ls Downloads/*.pdf` means “list anything in the Downloads folder that ends with ‘.pdf’,” whereas `ls Downloads/lab*` means “list anything in the Downloads folder that starts with ‘lab’.”

The table below summarizes the special symbols discussed here. Using these with the three command line navigation commands `pwd`, `ls`, and `cd`, you can locate and inspect any directory on your computer. You can also navigate to any folder and start IPython or Jupyter. Once you have found the proper directory, type `ipython` or `jupyter notebook` at the command prompt. You are ready to analyze that data!

Symbol	Refers To
.	the current directory
..	the directory that contains the current directory
~	your home directory
/	the root directory
*	wildcard character

⁴Section 4.1.2 mentioned the Windows format for pathnames, but Anaconda Prompt can use the format given here.

B.1.2 Creating, renaming, moving, and removing files

Suppose that you have a working script. You want to try out some new ideas for improving it, but you don't want to break a working version. One option is to create a duplicate and tinker with it, leaving the original intact. When you have worked out the bugs in your new version, you can archive or delete the original, or abandon the experimental version and go back to the original.⁵

The `touch` command is not in our table of essential commands, but it is useful for playing around at the command line. `touch original.py` will create an empty file called `original.py`. Type `ls` to see the new file in your current directory. To create a duplicate, type

```
cp original.py experiment.py
```

Type `ls` again to see the pair of files. You can change the name of the new file with the `move` command:

```
mv experiment.py next.py
```

Type `ls` again, and notice that `experiment.py` is gone, but `next.py` is now present. The `mv` command can also move a file from one directory to another.

To delete the original, type

```
rm original.py
```

or overwrite it with

```
mv next.py original.py
```

Type `ls` once more to see the current contents of the directory.

The command line offers a simple way to clean up after running a program that creates a lot of temporary files. For example, the `waves.py` script in Chapter 8 creates 100 image files for an animation. When you are done with this, you may want to reclaim the space on your hard drive. You can do this easily using `rm` and the wildcard character `*` discussed above: Navigate to the folder where these files were created and type `rm *_movie.jpg`. This will remove any files with names that end with `_movie.jpg` but will leave all other files in the directory untouched.

Most of the file creation in this book is done with Python. Deleting files using the command line is a good way to keep your file system organized. However, we remind you that these commands are unforgiving: `rm` does not preserve any copy in your Trash; `cp` and `mv` will destroy an existing file with the target name (without warning you), and so on. You should keep backups of your file system if you use command-line tools.

B.1.3 Creating and removing directories

When you start a new project or a new assignment, it is useful to keep all the files in one place, separate from everything else on your computer. Creating, moving, and removing directories is another set of tasks well suited to the command line.

If you don't already have a `scratch/` directory for temporary work, you can create one now. Navigate to your home directory, and then type `mkdir scratch` to create the directory. Type `ls` to see the new directory, and then `cd scratch` to move into it. Use `pwd` to confirm your new location. You can keep all of your Python work here, and you can use the same process to create folders for new assignments: for example, `mkdir assignment01`.

⁵ Section B.3 introduces more sophisticated version control tools.

The previous section described making a copy of a file to tinker with. You can do the same with an entire directory. First, create a new directory: `mkdir dir1`. Now, move into it (`cd dir1`) and create a few files (`touch fileA fileB fileC`). Move back up to the original directory (`cd ..`). To copy the directory and all of its contents, type

```
cp -r dir1 dir2
```

Typing `ls` will show that you now have two directories: `dir1` and `dir2`. You can view their contents: `ls dir1` and `ls dir2` should reveal identical file names in both. To remove the duplicate and all of its contents, type `rm -r dir2`. Use `ls` to verify the results of this command.

The option `-r` in these examples tells the command line to carry out the command (`cp` or `rm`) recursively—to do it to the given directory name and everything it finds inside. `rm -r` is a command you should use with care. Be sure that you really want to delete the folder and everything inside of it before you press <Return>.

B.1.4 Python and Conda

Most of the commands described so far have nothing specifically to do with Python. They just allow you to move around in your computer’s file system and to create and remove files or directories as needed. But our reason for discussing how to do these things is to make Python programming easier. You can do all of your Python programming from the command line if you like.

As mentioned earlier, you can start IPython, Jupyter, or Spyder directly from the command line. To start an IPython session, just type “`ipython`” at the command prompt and press <Return>. Python will have access to all of the files in the directory you were just working in. Typing `jupyter notebook` instead will launch Jupyter Notebook in your browser; again, you will see that it starts in the directory you were just in. This behavior is convenient if you need to access data files or scripts in that folder. (In contrast, Spyder will always start in its own default directory, regardless of your present working directory. You can adjust Spyder’s default directory through its Preferences menus.)

When you launch IPython, Jupyter, or Spyder from the command line, that application takes over the terminal window until you exit. If you wish to enter operating system commands, simply open a second window in your terminal app.

You can also run the Conda package manager from the command line to install and update Python packages and other programs.⁶ For example, to get the latest version of NumPy, you can type the following two commands:

```
conda update conda
conda update numpy
```

After each Conda command, you’ll be asked to confirm that you do want the available updates. You can even run Python scripts directly from the command line:

```
python my_script.py
```

Of course to run a script, you first need to write a script. And for that, you will need an application called a “text editor.”

⁶ Appendix A discusses Conda.

B.2 TEXT EDITORS

When working at the command line, you frequently need to edit plain text files, including Python scripts. For this purpose, you will need a text editor; a word-processing app such as Microsoft Word won't do. There are many choices. Choose one. Learn to use it. Learn to love it. Text editors often have a steep learning curve, but they tend to grow on you.

Almost all text editors have the ability to search and replace text. A good text editor for programming will have three additional, essential features.

- **Autoindent:** Your editor should have an *autoindent* feature that will ensure proper indentation and alignment in blocks of code.
- **Bracket Matching:** Your editor should identify unmatched brackets such as a “(” without a “)”, or an extra “].”
- **Syntax Highlighting:** This useful feature displays commands from Python and other programming languages in different colors and fonts. (The editor will use the file extension—for example, .py—to determine the language.) You can spot many coding errors at a glance with syntax highlighting.

Text editors with these features are available for every operating system. Vim and GNU Emacs are classic text editors included with most Unix and Linux systems and available for installation on Windows. They do everything, but they require some practice. GNU nano (“nano” for short) is a simple editor that still has all of the essential features (nano-editor.org). Atom is a modern open-source editor that runs on all operating systems (atom.io). Notepad++ is another free and open-source option on Windows (notepad-plus-plus.org).

If you are just getting started and don't already have a favorite text editor, you can install nano using the conda package manager.⁷

```
conda install --channel conda-forge nano
```

To edit a file, type `nano hello.py` at the command line. You should see a blank file with a description of useful commands at the bottom.⁸ Create a simple script. `print("Hello!")` will do. To save the file and exit, type `<Ctrl-X>`. Nano will ask if you want to save the changes you made. Confirm that you do, and accept its suggested location for saving the changes. Now run the file from the command line:

```
python hello.py
```

You have just created and executed a working Python program entirely from the command line!

Of the Big Three features mentioned above, nano's syntax highlighting and autoindentation are active by default. To match brackets, move the cursor over a bracket. Then, press `<Opt-]>` or `<Alt-]>` to jump to the matching bracket or display “No matching bracket.”

You can get a brief rundown of nano's commands by opening the editor and typing `<Ctrl-G>`, and you can learn more at nano-editor.org.

If you have launched IPython from the command line, then IPython's magic commands will assist you in editing files and running scripts. For example, `%edit my_script.py` will open `my_script.py` in your default text editor. Once you have created or edited the file, save it and quit the editor. IPython may automatically run your file. You can also run the script by typing `%run my_script.py`.

⁷ Alternatively, use `conda install --channel conda-forge/label/gcc7 nano`.

⁸ Nano's help text uses the common abbreviation `^X` to indicate `<Ctrl-X>` and so on. Nano takes over its terminal window until you exit the program. You can open a second window in your terminal if you need it.

You can set your default editor by modifying an IPython configuration file—from the command line! Navigate to your home directory, then move into a hidden folder:

```
cd ~/.ipython/profile_default
```

Use `ls` to see if you have a file named `ipython_config.py`. If not, create it with the command

```
ipython profile create
```

You can now edit this file to change your IPython preferences:

```
nano ipython_config.py
```

Find the following pair of lines:

```
## Set the editor used by IPython (default to $EDITOR/vi/notepad).  
#c.TerminalInteractiveShell.editor = 'vi'
```

Uncomment the second line (by deleting the `#`) and change `'vi'` to `'nano'` or your editor of choice. Save and close. The next time you type `%edit` in IPython, it will open your selected text editor.

B.3 VERSION CONTROL

You may have heard of GitHub. This is a website where programmers share their code and work together to make some amazing software. If you imagine the logistics of hundreds of people scattered across the globe all working on the same programming project, you might wonder how it is possible to coordinate their efforts and produce anything useful. Do they take turns editing programs? Do they share their work via email attachments? Actually, they use a **version control system**. It keeps track of the changes different programmers make and merges them together into a single working version of the project.

Git is a version control system. Rather, Git is *the* version control system. There are others, but Git is to version control as Google is to Internet searches. It is used by tens of millions of programmers, scientists, and engineers around the world to manage computer programming projects—everything from websites to the Linux kernel and artificial intelligence.

Git works by tracking changes in a **repository**. A repository is like a directory, but it has special folders and files to keep track of its history. Git can also synchronize repositories on your local computer with “remote” repositories stored on the Internet at sites like GitHub. If all you want to do is download files, you don’t need an account. But if you want to store your work on the Internet, you will need an account on a server site. GitHub (github.com) and Bitbucket (bitbucket.org) both offer free basic accounts. We recommend setting up an account now to try out all of the commands below; for concreteness, we will discuss GitHub.

You can do a lot with Git. This introduction will get you started with some of the most useful commands. Using Git is a great way to back up your work, share it with others, and collaborate with other programmers.

B.3.1 How Git works

Before we start using Git, we would like to explain two aspects of how it works: snapshots and local versus remote. This will help you understand some of the commands below and use the software more effectively.

effectively. We also describe how Git manages memory, but you can skip this on a first pass.

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Snapshots

It is important to understand the difference between the files you see in your folder and what Git actually keeps track of. Git only tracks the files you ask it to, when you ask it to. It does not track every modification you make. The first time you add a file to a repository, Git saves a **snapshot** of the file: a hidden, compressed version of the file. If you modify the file and save your changes, this has no effect on Git. Upon your request, though, Git will save another snapshot. Thus, Git stores the history of the repository as a series of snapshots of individual files. It can quickly reconstruct any snapshot of any file, or even merge different snapshots of the same file. Some of the commands that follow tell Git when to take a snapshot (`add`), which snapshots to store in the history (`commit`), and which collection of snapshots you wish to work with (`branch`). Thinking of this collection of snapshots may make some of the commands less mysterious.

Local versus remote

Git can help you track files on your computer, back them up online, share them with others, and copy a project to another computer. It does this by maintaining both **local** and **remote** repositories. A local repository is one that exists on the computer you are working at. A **remote** repository is one that exists online, often at a site like GitHub or Bitbucket. Two important commands discussed below are used to update your local repository: `add` and `commit`. These will only track the changes on the computer you are working on, with no effect anywhere else. If you want to back up your work online or update your local files with the latest versions from the online repository, you need to tell Git to transfer files between the local and remote repositories. The most common commands for this purpose are `push` and `pull`.

Changesets and file sizes

Keeping track of every change to every file in a repository sounds like it might use a lot of memory and take a long time to transfer files. This is generally not the case, thanks to Git's efficient compression algorithms and memory management.

First, the snapshots are compressed. Second, Git never stores a duplicate. The snapshot stores only the *content* of the file. Git can tell if two snapshots are really the same. Even if you have dozens of copies of the same file with different names in your folder, the Git repository only holds one compressed snapshot (plus a list of all the file names that share that content). Third, from time to time, Git compresses the already compressed snapshots for even more efficient storage and transfer of files. It stores the current compressed snapshot of each file and a compressed history of changes it can use to recover any previous snapshot. This compressed history contains “diffs” or “changesets”: Rather than storing two different versions of a file, Git stores the current version and the *changes* that must be made to the current version to recreate an earlier version. All of this makes Git a fast, flexible, and surprisingly efficient way to store the complete history of your project.

Git is most efficient when working with plain text files like Python scripts. Changes from one snapshot to the next are often minor: you add a new function to a module, add a few comments, and delete some unnecessary plotting commands. The changeset is minuscule, and storing the history of the repository as compressed changesets is quite efficient. However, for other types of files, a seemingly small change can have major effects. For example, a JPEG file stores an image in a compressed format. If you make a small change to the image you see on the screen, the compression algorithm could produce a completely different `.jpg` file. The changeset is almost as large as the file itself. Since there is no way to compress

The history of changes if many snapshots are completely different from the others, Git will compress the history.

the history of changes if every snapshot is completely different from the others, Git will essentially have

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to store every version of every image in the repository.

Many popular image formats have this property: JPEG, PNG, TIFF, GIF, or BMP. So do other popular file formats like PDFs, word processor documents (for example `.docx`), and spreadsheets (for example `.xlsx`). There are viable plain text alternatives for many applications: unformatted text, Markdown, L^AT_EX, or HTML instead of word processor formats; comma-separated or tab-separated values (`.csv` or `.tsv`) instead of spreadsheets; SVG and PostScript formats for images. Such files are often easier to create, load, and process with programs and scripts you write, too.

We recommend using Git for version control no matter what kind of files you work with. Git will faithfully track your project. Just be aware that repositories may contain some very large hidden files to track what seem like minor changes unless you are using plain text formats.

Overview

To summarize, a repository is a collection of snapshots of the files in a project. Repositories can be local or remote. Git works with any set of files, but it is ideal for plain text.

The remainder of this appendix will help you set up and start using Git. There are many different ways to use Git, but we focus on a minimal set of tools. We explain how to create local and remote repositories, how to track files, how to establish a connection between a local and a remote repository, and how to synchronize them. A typical session might go something like the following:

1. Get started. Use Git to create a new repository or to update your local repository with the latest changes to a remote repository.
2. Work on your project. Changes some files. Save your work.
3. Tell Git to track the new changes.
4. Repeat steps 2 and 3 as needed.
5. Use Git to upload the changes you made, from your local repository up to the remote repository.

Now let's get to know Git!

B.3.2 Installing and using Git

We recommend that you set up an account on github.com before you start. It is simple and free. You can then use your account as you work through the commands that follow. Many of the commands work fine without an online account, because they only affect your local repository. However, `clone`, `pull`, and `push` require an online repository, and `push` requires an online repository that you have permission to change (for example, one that you created).

Install Git

If you are using Linux or macOS, you already have Git installed on your computer. To confirm that, open a terminal and type the following at the prompt:

```
git --version
```

If you are using Windows, you will may need to install Git. You can download the software at Git for Windows (gitforwindows.org). Once you have installed the program, you should be able to run Git from the Anaconda Prompt. If that fails, you can use the Git Bash app (from the same website) instead.

Now you are ready to start using Git. Open a command line—terminal, shell, Anaconda Prompt, or the Git Bash app—and work through the examples that follow.

Note that Git has short abbreviations for many options used in the commands below to save typing. We use the long versions for clarity here, but you are likely to encounter the abbreviations elsewhere.

Set up Git

Before you start managing files with Git, you should specify some options to make working more convenient. Type the following commands at the command line to tell Git who you are and which editor you would like to use. (The default is `vi`. If you don't know what that is, you should change it now.)

```
git config --global user.name "<Your Name>"  
git config --global user.email "<username@myschool.edu>"  
git config --global core.editor "nano"
```

Create a local repository

Git tracks changes to files within a folder. If you want to start tracking changes to a collection of files on your computer, move into the directory that contains these files—or a new, empty directory—and type `git init`

This will create a hidden folder inside the current directory that stores its history. (The folder is called `.git`. You can look around in this directory, but do not change its contents or delete it.) You now have a local repository.

Create a new remote repository

You can use your account on GitHub to create a remote repository. Navigate to `github.com` and click on the “octocat” icon at the upper left. This will display your existing repositories, and it contains a button that will allow you to create a new repository. Click this button, give the repository a name, choose whether it will be public or private, and click “Create Repository.” You now have a remote repository. As long as it is empty, GitHub will describe several methods for putting files into it.

Copy an existing remote repository

Instead of creating an empty repository, you can make a copy of an existing project on GitHub. For example, you can **fork** the repository of code samples that appear in this book. Use the search bar on GitHub to find `dr-kinder/code-samples`. If you visit this repository—or any of the millions of other repositories on GitHub—you will see a button near the top right corner of the page that says “Fork”. Click on it. You now have your own remote copy of these files, to edit and experiment with as you see fit.

Forking is a feature of websites like GitHub and Bitbucket, not a feature of Git, but it is a common way to share code and collaborate. Once you have forked a project into your account, you can copy it to your local computer and start editing.

Copy a remote repository to a local repository

Sites like GitHub and Bitbucket are great for storing and sharing files, but to edit files and run programs, you will probably want to copy these files to your computer.⁹ Git provides a command to do exactly this: `clone`. To create a local copy of the remote repository of code samples on your own computer, open a terminal, navigate to the folder where you want the folder of code samples to be saved, and type

⁹ This may not always be the case. For example, Google Colab “is a Python development environment that runs in the browser using Google Cloud.” You can edit, run, save, and share Jupyter notebooks without storing any files on your computer. You can

even link it to your GitHub account. See colab.research.google.com.

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```
git clone https://github.com/dr-kinder/code-samples.git
```

(Replace “dr-kinder” with your own GitHub username to clone your own forked copy of the repository.) You will see a few messages from Git as it is cloning the directory onto your computer. When it is finished, you should see a new folder called `code-samples` that contains all of the code samples from this book.

If you attempt to clone a private repository, you may be asked for a password.

Link a local and a remote repository

You can now create a connection between a local repository (the files stored on your computer) and a remote repository (your files in the cloud). This step is only necessary if you created a new remote repository, or if you do not have permission to write to a repository you cloned. (Thus, you can skip this step if you forked the `code-samples` repository to your own account on GitHub, and then cloned the forked copy to your local computer.)

To establish a link, use the command line to navigate to your local repository, then type

```
git remote add origin https://github.com/username/my-repository.git
```

(Use your `username` and replace `my-repository` by the name of the new remote repository you created or cloned on GitHub.) You may be asked for GitHub credentials at this point. Git will remember them for future use.

`origin` is a standard nickname for a remote repository linked to a local repository. You can type “`origin`” instead of “`https://.../`” to save typing later. We can now share and synchronize files between the local repository on your computer and the remote repository on GitHub, all from the command line.

Create multiple versions of a project

You may want to keep the original code samples for reference, but tinker with them to learn how they work. Git allows you to create separate **branches** within a repository that can facilitate this. Each branch is an independent version of the project: Changes in one branch have no effect on other branches. However, since Git is tracking *all* of the changes to the repository, you can share changes between branches. This means you can always have a stable version of your project as you try out changes in separate branches.

If you are following along with the `code-samples` examples, move into the directory that you previously created via `git clone`. Make sure you are in the proper location. The following commands will not work properly unless you are in the `code-samples` directory. To see if you are in the correct location, type

```
git status
```

If there is a problem, you will see a message that begins with “`fatal: not a git repository.`” This means you are not in the correct directory. If things went well, you should see a few lines that begin with “`On branch master`.”

`master` is the default name for the main branch of a project. When you create a new repository, it is the *only* branch. You can create a new branch with the following command:

```
git branch archive
```

There is now an independent branch of the project named `archive` that contains the original, unmodified version of the code samples regardless of changes you make to those in the `master` branch.

Now, let's create a separate branch for **development** and "check out" this branch (like a book from the library).

```
git branch dev  
git checkout dev
```

This branch will track changes to the files without modifying either the `master` or `archive` branches. Changes will only affect this development version of the project, but you can import changes from other branches, too.

You can also add new branches to a remote repository. This makes it easy to share your work with others or pick up where you left off if you switch to another computer. To add your development branch to your online repository, use the following command:

```
git push --set-upstream origin dev
```

(You may be asked for your GitHub password.) `push` is a Git command that applies changes from a branch of your local repository into a branch of a remote repository. The `--set-upstream` option tells Git to remember this remote branch for future use with the `push` command. This time, the only change to the repository was creating a new branch. If you visit your site on GitHub, you should now see a new branch of your project. Next, we'll see what happens when you change the files in a branch of your project.

B.3.3 Tracking changes and synchronizing repositories

See what has changed

Edit a file in the folder or create a new one and type some text. Save your work, and quit the editor. Even though you have saved the modified file on your local computer, the changes are not yet part of the repository. To see changes that are not yet part of the repository, type

```
git status
```

You should see a message about one or several modified files.

Add snapshots to the local repository

When you are done making changes—or ready to save changes along the way—you must **add** and **commit** them to the repository for Git to keep track of them.

```
git add .  
git status  
git commit
```

The first command tells Git to take snapshots of all of the files you modified. (This operation is sometimes called "staging.") The period after `add` means "everything that has been created or modified since last time." You can provide an individual filename or a list of filenames instead. The second command shows the changes effected by the first. The third command makes the new snapshots a permanent addition to the repository.

`git commit` will open a new, empty file in a text editor. This is your opportunity to describe the

changes that you are about to save to the repository. You should summarize or explain the changes with a short message on the first line. If you wish to write more, leave a blank line and then include a more detailed description. Save and quit the editor. Git has now made the changes part of the repository. You can see the effect by typing `git status` again.

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Export changes

You have updated your local repository, but not your remote repository. It is time to push these changes from your computer to GitHub.

```
git push
```

That's it! All the changes you committed in this branch of your local repository are now part of the corresponding branch of the remote repository. You can clone the online repository to a different computer and pick up where you left off. You can rest easy knowing you have a backup copy of your work. Use `add`, `commit`, and `push` frequently.

`push` can export changes from any branch of one repository to any branch of another. Here, we took advantage of the `--set-upstream` option we used earlier, when creating the remote branch. Git will automatically push changes from this local branch to that remote branch, unless you explicitly instruct it to do otherwise.

Merge branches

When you have finished tinkering, debugging, and editing in the development branch, you can use the `merge` command to incorporate your changes into another branch. Check out the branch you want to update, and then specify the branch you wish to import changes from with the `merge` command.

```
git checkout master  
git merge dev
```

Now all of the changes you made in the `dev` branch are part of the `master` branch, too. This merge has no effect on `dev`. You should probably push the changes to your remote repository, too. You can use branching and merging to work on several different tasks within a project at the same time, all while keeping a working version available until the new changes are ready. Different developers can work in different branches on the same project, too, and pool their changes in the `master` branch.

Import updates

Suppose that you make changes on one computer and push them to your remote repository. If you start working on a different computer, you will want to update its local repository before you start making new changes. Or, if you are collaborating with others and everyone is pushing their changes to the same remote repository, you will want to update your local repository with the latest version of the project from time to time. Use Git to *pull* changes from a remote repository into a local repository.

Commit your local changes before you pull.

If you do not commit your latest changes, they may be overwritten and lost.

```
git add .  
git commit  
git pull origin master
```

`pull` will import new files and changes in the `master` branch of the remote repository into the current branch of your local repository, and then merge these into your current version of the project.¹⁰

¹⁰If you cloned a repository that has still yet to update into your local repository, now, if you don't have

If you cloned a repository, you can still `pull` updates into your local repository, even if you don't have permission to push to that remote repository.

¹⁰ `pull` combines two separate Git commands: `fetch`, which copies the changes into your repository without implementing them, and `merge`, which attempts to implement all of the changes.

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Ignore files

You don't have to track every file in a folder. You can choose not to add certain files, or you can give Git explicit instructions to ignore them. One approach is to create a hidden file called `.gitignore` and maintain a list of the files Git should ignore. Open and edit it like any other plain text file. (See Section B.2.) For instance, a `.gitignore` file might have the following lines:

```
figure1.pdf  
figure2*  
*.jpg
```

(Git understands the wildcard syntax when processing a `.gitignore` file. See Section B.1.1.)

This file instructs Git to ignore `figure1.pdf` (but no other PDFs), any file whose name starts with `figure2` (such as `figure2.jpg`, `figure2.png`, and `figure2.pdf`), and *all* JPEG files (`figure1.jpg`, `figure2.jpg`, and so on.). You can still add files that would otherwise be ignored:

```
git add figure2.jpg
```

If a file is *already* part of the repository, adding its name to a `.gitignore` file will cause Git to stop tracking changes, but it will not remove the file or its past history from the repository.

Dealing with conflicts

Git is quite proficient in merging files. Even if multiple authors have made changes to the same file, Git can usually weave them together into a single coherent version automatically. However, if two authors change the same line of a file, or if you make different changes to the same line of a file in two different branches of a repository, Git does not know which one to keep. Instead of guessing, Git will flag the conflict, alert you to it, and suspend the `merge` or `pull` operation until you resolve the conflict. Git will issue a warning like this:

```
CONFLICT (content): Merge conflict in file.txt
```

If you open the file, you can see where Git ran into trouble. It indicates the conflict by inserting extra lines into the file:

```
<<<<< HEAD  
You can't change this line.  
=====  
I can change this line.  
>>>>> dev
```

In this example, the file in the current branch (denoted by `HEAD`) contains a different sentence than the `dev` branch at the same location in `file.txt`. The two conflicting versions are separated by equal signs. To resolve the conflict, choose one sentence and delete the other, along with the extra lines that start with `<<`, `>>`, and `==`. Once you have resolved all conflicts in this way, `add` and `commit` the changes to complete the merge.¹¹

B.3.4 Summary of useful workflows

There are a few common tasks that combine multiple Git commands. We collect them here to help you get started.

¹¹ If you run into this problem often, you may want to look into a “merge tool” like Meld (meldmerge.org) or KDiff3 (kdiff3.sourceforge.net).

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Create a new project:

```
mkdir new-project
cd new-project
git init
...
[edit files and save]
...
git add .
git commit
```

You may want to link to a remote repository and push your changes at this point.

Clone a project. First, fork the project into your account on GitHub or Bitbucket. Then:

```
git clone https://github.com/username/new-project.git
cd new-project
git branch archive
git branch dev
git checkout dev
git push --set-upstream origin dev
```

Track and back up your work:

```
...
[edit files and save]
...
git add .
git commit
git push
...
[make more changes]
...
git add .
git commit
git push
```

Always do a final push before you quit working on a project, especially if you plan to resume your work on a different computer or share with a collaborator.

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Download updates to a project:

```
git add .
git commit
git pull origin master
```

Branching, merging, pushing, and pulling allow many people to work on many different tasks within a single project at the same time. These processes are integral to the organized chaos that is open-source programming. Now you can join in!

B.3.5 Troubleshooting

It's easy to make mistakes with Git, and the program is not user friendly for beginners. But it is powerful. You are unlikely to make any mistakes that cannot be fixed, especially if you commit and push your changes frequently. However, it may take a while to figure out how to fix problems. The following situations commonly arise when learning to use Git. We offer some black box commands that might help get you back on track.

I tried to commit my changes, but nothing happened. Is Git broken?

No. Type

```
git status
```

If you see a message that includes "not a git repository," it means you are in the wrong directory or that you never created a repository with `git init`. If you instead see a message about "untracked files" or "changes not staged for commit," that means you did not instruct Git to track your changes. Use

```
git add <filename>
```

to add a specific file, or

```
git add .
```

to add all of the untracked changes.

Oh, no! I messed up a file and saved the changes. Can I go back to an earlier version?

Yes. If you only need to undo changes to one file, use `checkout`:

```
git checkout -- <filename>
```

This will restore the file to its state at the last commit. You can revert your entire repository to its state at your last commit if you need to:

```
git reset --hard
```

Note that this will revert **all files** to their previous state, and you cannot undo this action.

Oh, no! I accidentally deleted all of my files. Does that mean I have to start over?

No. Just revert to the previous version of your repository, as described in the previous question. All your files will be back.

But I committed after I deleted the files. Does that mean I have to start over?

No. You can roll back to an earlier commit. You need to look up a unique label for an earlier commit. To see a record of all of your previous commits, use the `log` command.


```
git log --oneline
```

This will display a list of all the changes you have committed, along with a “commit ID”—a unique hexadecimal (base-16) identifier—and the comments you wrote. (Write good comments!) If you want to revert all of your files to a previous version, use the following:

```
git revert --no-commit <commit ID>..HEAD  
git commit
```

If you only want to revert a specific file, use `checkout` to get the version of the file you need:

```
git checkout <commit ID> -- <filename>
```

Oh, no! I forgot to check out my development branch and committed changes to the master branch. I don't want to overwrite that branch when I try to push. Do I have to start over?

No. You can pull from one local branch to another. First, check out the local branch you wanted to use; then, merge the changes from the branch you unintentionally committed them to.

```
git checkout dev  
git merge master  
git commit
```

Now you should be able to push your changes to your remote development branch of the project. To undo the changes you made to the master branch, use the `revert` command, as described in the previous question. Only do this after you commit your changes in the development branch. They will be gone from the master branch after this command.

Oh, no! I changed some of my files, forgot to commit the changes, and then I accidentally deleted them. Does this mean I have to start over?

Yes. You should commit your changes frequently. Git is version control software, not a time machine.

More information

We have only scratched the surface of what Git can do. If you want a more thorough introduction, see Chacon & Straub, 2014 (git-scm.com/book). For a hands-on introduction, visit the GitHub Learning Lab (lab.github.com) or the tutorials at Bitbucket (atlassian.com/git/tutorials). You can also learn as you go with Google and Stack Overflow.

B.4 CONCLUSION

This appendix is a lot to digest, and if you have never worked at the command line, it may seem intimidating. However, learning these tools can improve your programming experience and make you a better programmer. Some things are easier at the command line, and version control is a marvelous invention. A good text editor ties it all together.

APPENDIX C

Jupyter Notebooks

There are many ways to write and run Python code. You could simply write scripts with a text editor and run them from your operating system's command line. But many "front end" systems have been developed to make coding easier. One of these is the Spyder integrated development environment (IDE) described throughout the main text. This appendix describes another popular choice: the Jupyter Notebooks system.¹

Like a physical notebook, a Jupyter notebook is a record of your work. It is an interactive document that is displayed in a web browser. However, all files are stored on your own computer, not on a web server. Each notebook consists of a series of **cells**. Cells may contain either executable code or formatted text. This means you can include presentation-quality documentation alongside your source code. Jupyter notebooks are also easy to share. Another person can open your notebook and run or modify the code while reading your nicely formatted commentary.

Let's look at a Jupyter notebook now.

C.1 GETTING STARTED

Although you view Jupyter notebooks in your web browser, the notebook software must first be installed on your computer. This happens automatically when you install the Anaconda distribution. It can also be done separately with the `conda` package manager. (See Appendix A.)

C.1.1 Launch Jupyter Notebooks

Once you have installed the software, you can launch a Jupyter Notebook session from your operating system's command line with the command

```
jupyter notebook
```

You can also use the Anaconda Navigator application to launch the Jupyter Notebook application. (You may get a message like

```
Copy/paste this URL into your browser when you connect for the first time,  
to login with a token: http://localhost:8888/?token=XXXX
```

Do as you are instructed to get started.)

A window will open in your default web browser with "Jupyter" at the top left corner (Figure C.1).² The first of three panes is selected by default. This is a file browser that you can use to find and open an

¹ "Jupyter Notebooks" is an extension of "IPython Notebooks" that will eventually be subsumed into "JupyterLab." The interface is similar to the notebook environment of *Mathematica*, Maple, and Sage.

² **T2**Starting Jupyter launches several "background processes" that you do not see. These processes communicate with your web browser, which communicates with you. Both Jupyter and your notebooks work even if you are offline, because they are located on your computer. The web browser simply provides a convenient and familiar interface for human users.



Figure C.1: The Jupyter Notebook file browser.

existing notebook or create new notebooks. If you open a Jupyter notebook from your operating system’s command line or the Anaconda Prompt, the file browser will start in the directory from which you issued the command.³ If you launch from Anaconda Navigator, the file browser will start in your home directory.

C.1.2 Open a notebook

If you don’t have any notebooks yet, use the file browser and navigate to a folder where you’d like to create one. (You can create a new folder with the **[NEW]** button at the top right of the file browser. Check the box next to the new folder and then click **[RENAME]** to rename it.) Next, click on the **[NEW]** button and select “Python 3” from the dropdown menu. A new browser tab will open with a Jupyter notebook that contains a single empty code cell called `In []` (Figure C.2). Enter `1+1` and then click the **[Run]** button or hit `<Shift-Return>` on your keyboard. You have just run your first code in a Jupyter notebook!

You can save your notebook with the **[Save]** button, the Jupyter menu item **File>Save**, or the shortcut `<Cmd-S>`. (Use the menu inside the Jupyter window — not your browser’s **File>Save** menu item.) You can now find it in your computer’s file system (probably named `Untitled.ipynb`). If you would like to rename a file, you can simply click on the name of the notebook at the top of the page, or you can use the menu option **File>Rename....**

You can also open existing notebooks saved on your computer. For example, you can view a sample notebook at the blog that accompanies this book.⁴ It illustrates many features of Jupyter notebooks. You can download the notebook, open it, modify it, and play with your own copy. Many other notebooks are also available for download from the web.

C.1.3 Multiple notebooks

You can run multiple notebooks at the same time. Return to the file browser tab, create a second new notebook, and perform some other calculations. You now have two independent notebook sessions. Each one has its own state, including the values of all variables. No information is shared between notebooks. You can keep track of all open notebooks in the file browser tab: just select the **[RUNNING]** pane. From this pane, you can selectively terminate any of your running notebooks.

³ See Section B.1.1

⁴ physicalmodelingwithpython.blogspot.com/2016/01/jupyter-notebooks.html.

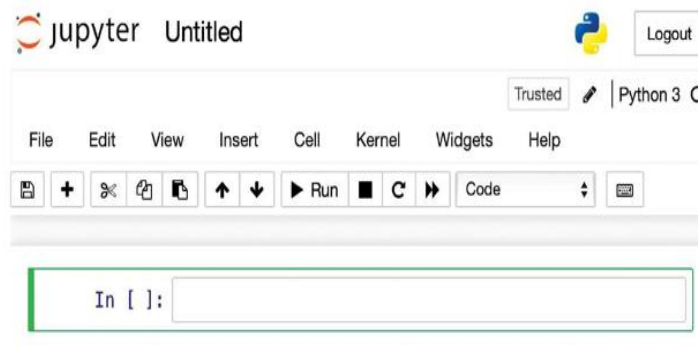


Figure C.2: A new Jupyter notebook. The green border on the selected cell indicates that Jupyter is in input mode.

C.1.4 Quitting Jupyter

When you are finished computing and editing, you will need to be sure you terminate all of the processes Jupyter is running on your computer. Simply closing the tabs in your web browser will not do the trick.

First, shut down each notebook. From within a notebook, use the menu option `File>Close` and `Halt`. This will close the notebook and shut down the Python process that was running in the background. Alternatively, you can use the file browser. Select the “Running” pane to see a list of all active notebooks. There will be a `[SHUTDOWN]` button next to each. Use it to shut down all active notebooks.

Next, you need to terminate the Jupyter Notebooks application itself. The simplest way is to use the `[QUIT]` button (Figure C.1). Alternatively, locate the “terminal window” associated with the application. If you launched the Jupyter Notebooks application from the Anaconda Navigator, you will need to find an application window that displays a series of messages from `NotebookApp`:

```
[I 03:14:04.679 NotebookApp] ...
[W 03:17:26.088 NotebookApp] ...
...
```

If you started Jupyter Notebooks from your operating system’s command line, this will be the terminal window where you entered the startup command. Click on this application. Close the Jupyter Notebooks application once and for all by typing `<Ctrl-C>`. The program will ask for confirmation. Press `<y>` then `<Return>` to confirm. The program is now closed.

The remainder of this appendix provides a brief overview of some useful features of Jupyter notebooks. However, the best way to learn how to use Jupyter notebooks is to create your own and explore those created by others.

C.1.5 `T2`Setting the default directory

It is good practice to keep all of your work in one location. You can set the default starting folder for Jupyter notebook sessions. (This is analogous to choosing the “current working directory” in Spyder.) However, this is not currently possible from within Jupyter. You must use your operating system’s command line. At the command line, type

```
jupyter notebook --generate-config
```


This will generate a configuration file called `jupyter_notebook_config.py`. This is simply a Python file that sets a lot of options every time Jupyter starts up. It will be located in a hidden folder called `.jupyter` in your home directory. Open this file in a text editor. Search for the lines

```
## The directory to use for notebooks and kernels.  
#c.NotebookApp.notebook_dir = ''
```

Uncomment the second line and replace the empty string with the name of the folder where you want Jupyter to run, such as `"/Users/username/scratch/jupyter"`. Save and close. The next time you start Jupyter, it will open in this folder.

C.2 CELLS

When you created a new, empty notebook, it consisted of a single cell. Every Jupyter notebook is composed of one or more cells. Each cell may be one of the following three types:

- Executable Python code (or code in more than 100 other programming languages), possibly accompanied by output from that code;
- Text formatted with the Markdown system;
- Plain text.

You can select any cell and then insert a new one before or after it by using the `Insert` menu.

C.2.1 Code cells

Code cells are those in which you enter Python commands. A single code cell may contain multiple Python statements. If the code generates output, it will appear below the input after an `Out [N]:` prompt.

Help is available in code cells. Type `np.power?` in a cell and run it: A window pops up with documentation. `help(np.power)` displays the same information within the current cell.

A series of code cells resembles a Python script, but Python will not necessarily carry out commands in the order you entered them. When you create a new notebook or open an existing one, Jupyter launches an IPython **kernel** in the background. This is the program that executes all of the Python code during your session.⁵ You enter your code in code cells, and you can run these in any order you choose. Or you can select a group of cells and run them all in sequence by using the `Cell` menu.

The notebook informs you of the order in which cells were evaluated. When you created an empty notebook, the cell was labeled `In []`. When you asked Jupyter to evaluate `1+1`, the name of the code cell changed to `In [1]` to indicate that this was the first cell you evaluated in that session. When you have many cells, they can be run in any order you wish. The cell name indicates the sequence number of the most recent evaluation of that cell.⁶

Each notebook remembers its state as you execute various cells. The effect of executing a cell on Python's state persists even if later you delete that cell. To start fresh, you can restart the kernel. Use one of the restart options available from the `Kernel` menu at the top of the page. There is also a  button for this purpose.

⁵ Jupyter Notebooks and the Spyder IDE are built on top of the same IPython interpreter. All the Python code and IPython magic commands in this book can be run in code cells.

⁶ When a cell takes a long time to run, its name temporarily changes to `In [*]` to indicate that it is still being processed.

After you terminate or restart a kernel, all of your output remains visible in the code cells, even though the state of the kernel has been lost. That's good if you want to save the notebook or share it with another reader. However, this behavior can be confusing if you want to rerun your notebook from scratch. You can easily get confused about which output came from the latest run and which came from the previous one. For this situation, there is a convenient Jupyter menu item called `Kernel>Restart & Clear Output`.

C.2.2 Graphics

If the code in a cell generates a figure, that will appear in the output region of the cell by default. (You may need to add a `plt.show()` command to render graphics.)

If you prefer an interactive graphic instead of the default “inline” behavior, include the following “magic” command immediately after importing PyPlot:

```
%matplotlib notebook    # Create interactive graphics.
```

(This should eliminate the need for any `plt.show()` commands.) New figures have a “power button” on the upper-right corner of the interactive plot window. Do all of your adjustments and explorations before you click on it. Once you click on it, the figure will be frozen in its current state until you run the cell to generate it again.

If you want to switch back to inline graphics, use the magic command

```
%matplotlib inline      # Display static output in the cell.
```

C.2.3 Markdown cells

New cells are code cells by default, but you may wish to add some explanation, commentary, or reflection. You could use Python comments within code cells, but Jupyter notebooks offer a more flexible method for entering text. Select one or more cells and use the `Cell>Cell Type` menu to change them to Markdown mode. Now, you can type text in the cell that will not be executed. Your typing shows up as plain text as you type. When you are finished, evaluate the cell with the button, as you would a code cell. The plain text input is replaced by more aesthetically pleasing output. If you want to change what you have typed, double-click on the cell. You will see the plain text that you entered, and you can begin editing.

You can format text using the Markdown language. Cells can even contain mathematical expressions rendered with MathJax.⁷ A description of these features is beyond the scope of this appendix, but the sample notebook demonstrates some of the possibilities. A web search for “jupyter notebook gallery” provides many examples.

C.2.4 Edit mode and command mode

A Jupyter notebook has two primary modes of operation. In command mode, the keystrokes carry out operations within the notebook: saving text, evaluating code cells, inserting or deleting cells, and so on. In edit mode, typing produces text within a cell.

You can enter edit mode by double-clicking on any cell. If a cell is highlighted, you can also enter edit mode simply by pressing `<Return>`. (Keep in mind, however, that editing a cell has no effect on

⁷ MathJax allows you to use most \TeX and some \LaTeX commands to produce beautiful mathematical output in web pages. See docs.mathjax.org.

the kernel’s state until you actually run the cell.) You can leave edit mode and enter command mode at any time by clicking outside of a cell, or by pressing the `<Esc>` key. Now many common operations can be carried out with a single keystroke. For example, `<X>` will delete the current cell. If you accidentally press `<X>` while in command mode, you can undo the operation with `<Z>`. Another useful shortcut is `<S>`, which saves your notebook. Type `<H>` in command mode or see `Help>Keyboard Shortcuts` for the complete list. Learning a few keyboard shortcuts can make using Jupyter notebooks a lot easier and more efficient.

C.3 SHARING

You can share a notebook file with anyone who also runs Jupyter just by sending them the `.ipynb` file. You can also export a static version in several other formats using the Jupyter menu item `File > Download as`. An HTML version of your notebook requires nothing but a web browser to view it. If you have the free `LATEX` software installed on your system, you can also convert the notebook to a PDF file. Finally, you can share your notebooks over the web.

If you save a notebook to Dropbox, you can share a link to the notebook with others. They can go to the Jupyter notebook viewer (`nbviewer.jupyter.org`), enter the link, and view your notebook—even if they do not have Python or Jupyter Notebooks installed on their computer. The code repository `github.com` even has a notebook viewer built into the website. If you upload your notebook there, anyone can view it, download it, or use a link to the file to view it in a notebook viewer. Another powerful collaboration system is the Google Colaboratory: `colab.research.google.com/notebooks/welcome.ipynb`.

You can use `File > Download as .py` to export your notebook to a standalone Python script. Everything but code cells will be converted to Python comments.

C.4 MORE DETAILS

The information in this appendix should be enough to get you started with Jupyter Notebooks. The notebook `Introduction.ipynb` available on the blog that accompanies this book demonstrates more techniques. For more information, have a look at `jupyter-notebook.readthedocs.io`. That site contains information on Markdown, MathJax, other programming languages, and many other topics. Much of this information is even available in Jupyter notebooks.

C.5 PROS AND CONS

Jupyter notebooks allow you to *do* your work, document your work, and see the results in a single file. This makes them very useful for exploring new ideas, solving problems, and collaborating. Jupyter notebooks are also useful for presenting your work. The notebook eliminates the often painful step of including code and graphics in a formatted document. (Try creating a similar document with a word processor, and you will see the advantages of Jupyter notebooks!) Jupyter notebooks are also useful in teaching. A teacher can use notebooks for demonstrations, in-class collaborative work, and assignments. A partially completed notebook can be a useful starting point for a student project.

Jupyter notebooks do many things very well. However, no tool is perfect for every job. The browser interface places another layer of software between the programmer and the machine, and performance can suffer. Complex graphics can overwhelm a browser at times. Many scientists and programmers prefer an IDE such as Spyder for writing and debugging large programs—especially those involving multiple files. It can be difficult to transform a notebook into a polished piece of software.

Give Jupyter notebooks a try. You may find them to be the perfect tool for some of your work.

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APPENDIX D

Errors and Error Messages

Now would be a good time to start making errors. Whenever you learn a new feature, you should try to make as many errors as possible, as soon as possible. When you make deliberate errors, you get to see what the error messages look like. Later, when you make accidental errors, you will know what the messages mean.

— Allen B. Downey

Everyone makes mistakes. When Python detects something wrong, it will halt a program and display an error message. This process is called “raising an exception.”

Of course, Python doesn’t really know what you are trying to accomplish. (At least not yet....) If you enter something syntactically correct, but not what you need to solve your problem, Python can’t point that out to you. Hence the importance of testing your code and debugging it. Even when Python does detect something wrong, it may not be able to tell you what you need to fix.

This appendix makes no attempt to catalog all of Python’s error messages. Instead, we show a few common errors and decode them. As you learn about how Python interprets your code, you’ll gain insight into these messages.

D.1 PYTHON ERRORS IN GENERAL

Before getting down to specifics, let’s take a look at Python’s general method for handling errors. Follow the advice at the beginning of this chapter and make a few errors. First, type `%reset` to erase all variables and modules. Then, type the following commands at the command prompt as written. (Feel free to make your own errors as well.)

```
1/0
import numpy as np
abs(-3)
a = [1,2,3]; a[3]
5 b = [1,2,3]
b[0]
```

You should get a different error message from each command. Notice that each error has a name—`ZeroDivisionError`, `TypeError`, `SyntaxError`, and so on—followed by a message. When Python detects an error, it raises an exception. You can raise an exception manually, and even provide your own message. Try these commands:

```
raise TypeError
raise ZeroDivisionError("You should not divide by zero!")
```


Python has classified many common errors. Thus, when it recognizes an error, it tries to describe what went wrong and provide additional messages supplied by programmers. These messages may not seem very informative at first, but compared with the “segmentation fault” of lower-level programming languages, they are quite helpful.

Python not only allows programs to raise exceptions; it also allows you to catch them before they terminate your program. To see the difference, try the following two methods of dividing by zero:

```
import numpy as np
np.divide(1, 0)
1/0
```

The programmers who created NumPy decided that sometimes it is OK to divide by zero, because some functions do have singularities. So they downgraded the exception from **ZeroDivisionError** to **RuntimeWarning**, thus allowing some singular functions to return a value.¹ This behavior allows NumPy to process an entire array and return a result, even if some of the entries involve operations that are not well defined mathematically. In contrast, basic Python stops execution and alerts the user.

You can override the default behavior by using Python’s protocol for handling exceptions. A function call may raise an exception, but you can catch it before it halts the entire program and do something else. Try typing

```
try:
    1/0
except ZeroDivisionError:
    print("1/0 -> infinity")
```

This syntax tells Python to “try” running the code in the first indented block. If there is an error, Python will inspect it. If the error is anything other than a **ZeroDivisionError**, Python will halt the program and print an error message. However, if it finds a **ZeroDivisionError**, it will run the code in the second indented block and proceed with no further warnings. This is an example of *exception handling*. It is a useful technique you can use in your own programs.

You do not need to master exceptions and exception handling at this point. The takeaway message from this section is that Python will try to alert you to the nature of the error by raising an exception, but not every program will handle the same errors in the same way.

Now let’s take a look at some common errors.

D.2 SOME COMMON ERRORS

- **SyntaxError** – This is the most common error for beginning programmers. It usually means you typed a command incorrectly. To generate a syntax error, try the following:

```
abs -3
```

It may be obvious to a human reader what you mean here, but Python is not a human reader. It knows of a function called **abs**, but this function should be called with a single argument in parentheses. You did not call this function using the proper syntax, so Python raises a **SyntaxError**.

¹ Some IDEs will suppress the **RuntimeWarning** and simply return the value **np.inf** without complaining.

Sometimes Python is able to pinpoint exactly where the error occurred. Suppose that you used a single equals sign when you wanted to test for equality:

```
if q = 3:
    print('yes')
else:
    print('no')
```

Python replies with

```
File "<ipython-input-87-19c154aec9ce>", line 1
  if q = 3:
          ^
SyntaxError: invalid syntax
```

Here the caret (“^”) pinpoints where Python’s scanner had arrived when it detected something wrong. A syntax error may actually be located in a line *preceding* the one that is flagged. Python waits until it is “sure” there is an error before issuing the exception. If you find that a flagged line looks correct, work backward to see if there is an error on an earlier line.

Unmatched parentheses are a common source of syntax errors. For example, try the following:

```
x = -3
print(abs(x)))
```

Python will also raise a **SyntaxError** if it reaches the end of a script while still looking for arguments to a function, closing parentheses, or something similar:

```
x = -3
print(abs(x))
```

If you type this in a script and then attempt to run it, you will get a **SyntaxError**. However, if you type these commands at the command prompt, IPython won’t allow you to commit the error. Every time you hit <Return>, the cursor moves down a line and nothing else happens. Just close the parentheses, hit <Return>, and Python will execute the command.

- **ImportError** – Python raises this exception when it cannot find a module you are attempting to import or when it cannot find the function or submodule you are attempting to import from a valid module. Most often, this occurs when you type the name of the module or function incorrectly. (Remember, Python names are case sensitive.) Each of these lines raises an exception:

```
import NumPy
import nump as np
from numpy import stddev
```

If you are certain that a module exists and you are spelling its name correctly, then you may need to install the module on your computer, perhaps by using the `conda` tool. (See Appendix A.) If you have installed the module, you may need to move it to a different directory or modify your computer’s `PYTHONPATH` environment variable in order for Python to find it.

- **AttributeError** – Python raises this exception when you ask for a data field or method from an object that does not possess it:

```
np.atan(3)  
np.cosine(3)
```

This is usually due to spelling the name of a data field or method incorrectly, or misremembering its abbreviation. (NumPy's names for the desired functions are `np.arctan` and `np.cos`.)

- **NameError** – Python raises this exception when you ask for a variable, function, or module that does not exist. This can result from misspelling the name of an existing variable, function, or module. It can also happen when you forget to import a module or forget to define a variable before using it.

```
%reset  
zlist  
np.cos(3)
```

A puzzling instance of this error can arise when you define a variable in a script or at the IPython console, and then attempt to use that variable in another script. Even though you can see the variable in the Variable Explorer and use it in the console, other scripts cannot access this variable. That is because IPython runs each script in its own private **namespace** and only brings variables, functions, and objects created by the script into your IPython session *after* the program finishes—or crashes. (See Appendix F for more about namespaces.) Writing scripts that do not define or import the variables and functions they use is bad coding practice. If you need to use the results of one script inside another, import the first script as a module within the second script.

- **IndexError** – Python raises this exception when you provide an index that lies outside the range of a list or array:

```
x = [1, 2, 3]  
x[3]
```

You may make this error a few times before you get used to Python's system of numbering elements starting at 0 rather than 1.

- **TypeError** – Python raises this exception when you call a function with the wrong type of argument. This can occur if you asked for user input and forgot to convert the string to a number before performing a mathematical operation. It can also occur if you combine two objects in a way Python cannot interpret:

```
abs('-3')  
x = [1, 2, 3]    # not an error  
x[1.5]  
2 + x
```

- **AssertionError** – Python raises this exception when an assertion statement is violated. It may be the most helpful error message of all, because you ask Python to alert you to the problem.

```
x = [1, 2, 3, 4]  
y = [1, 4, 9, 16, 25]  
assert len(x) == len(y), "Lists must be same length!"
```

Python has more than 60 built-in exceptions and warnings, each for a unique type of error. See how many you can generate!

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APPENDIX E

Python 2 versus Python 3

There are two major versions of Python in wide use: Python 2.7 and Python 3. In 2008, version 3.0 of Python was released. This version was the first to break *backwards compatibility*, meaning that code written in earlier versions of Python was not guaranteed to run in version 3.0 and later. Python 2 is no longer under active development. The last official release was on January 1, 2020. However, just because a language is not under active development does not mean it will disappear.¹

In this tutorial, we have chosen Python 3 when a choice was required. If you are already familiar with Python 2 and do not want to change, or your advisor insists that you use Python 2, have no fear. All the modules we describe are available in both Python 2 and Python 3, and we have taken care to write code that will work in either version whenever possible. There are only three instances where accommodating both versions was not practical: division, the `print` command, and user input.

Fortunately, there is a special module called `__future__` that makes many features of Python 3 available within Python 2. This includes both division and the `print()` function. To run all of the code samples in this tutorial in Python 2.7, you only need to add the following two lines at the top of each script and execute them at the beginning of each interactive session:

```
from __future__ import division, print_function  
input = raw_input
```

To understand why, read on. To try out code samples, you can return to the main text now.

E.1 DIVISION

In Python 2, division of two integers returns the quotient and ignores the remainder. Thus, the result is always an integer, even if the integers are not divisible. In Python 3, division of two integers instead returns a floating-point number if they are not divisible.

```
1/2 == 0          # True in Python 2; False in Python 3  
1/2 == 0.5       # False in Python 2; True in Python 3
```

The second option is usually what we want in numerical computation. Users of Python 2 can get this behavior from the `__future__` module:

```
from __future__ import division
```

Integer division is still available in both versions of Python:

```
3//2 == 1          # True in Python 2; True in Python 3
```

¹ No new features have been added to TeX since 1984; FORTRAN 77 and COBOL are still in wide use.

E.2 PRINT COMMAND

The `print` command behaves differently in Python 2 and Python 3, though this will not have a significant effect on the code samples in this tutorial.

In Python 2, `print` is a statement, like `assert`, `for`, `while`, and others. The command

```
print "Hello, world!"
```

will do what you expect in Python 2, but in Python 3 it will raise a `SyntaxError`. That's because `print` is a *function* in Python 3. It requires arguments enclosed in parentheses, like every other function.

In this tutorial, all print commands have the form required by Python 3:

```
print("Hello, world!")
```

This particular statement will also work in Python 2. However, if you use Python 2, be aware that it will interpret a series of arguments in parentheses as a tuple. Thus,

```
print('x', 'y', 'z')
```

will produce different output in Python 2 and 3.

To use the Python 3 `print` function in Python 2, import it from the `__future__` module:

```
from __future__ import print_function
```

There are two caveats. First, you cannot have both variants of the `print` command at the same time.

```
print "Hello, world!"
```

will now result in a `SyntaxError`. Second, importing from the `__future__` module is irreversible. You must restart Python to revert to standard Python 2 behavior.

E.3 USER INPUT

Python 2 has two statements for obtaining input from the user: `input()` and `raw_input()`. In contrast, Python 3 has only one: `input()`. However, Python 3's `input()` function behaves like Python 2's `raw_input()` function. That means code samples in this tutorial that use `input()` may not work properly in Python 2.

What is the difference between the two? In Python 2, `raw_input()` returns a string, whereas `input()` attempts to evaluate whatever the user types as a Python statement. Suppose a script has the following statements:

```
input("Type 33/3 and hit <Return>: ")
raw_input("Type 33/3 and hit <Return>: ") # Raises NameError in Python 3
```

In Python 2, if the user follows instructions, the first statement will return the integer 11.² The second statement will return the string '`33/3`'. In Python 3, the first statement will return the string '`33/3`', while the second statement will result in an error, because there is no function named `raw_input()`.

The exercises in this tutorial assume that `input()` returns a string. They may run without modification, but to run the code samples as intended in Python 2, you will need to either replace every instance of `input()` with `raw_input()`, or redefine the `input` function:

² Users who do *not* follow instructions are the primary reason this function was eliminated in Python 3.


```
input = raw_input
```

The `input` function is not available in the `__future__` module at this time.

E.4 MORE ASSISTANCE

A script named `2to3` automatically generates Python 3 code from programs written using Python 2. Though it is not necessary for this tutorial, `2to3` may be useful in making your other projects compatible with Python 3. See docs.python.org/2/library/2to3.html. Another option is the `six` module whose stated purpose is “to support codebases that work on both Python 2 and 3 without modification”: six.readthedocs.io.

It is also possible to install and run multiple versions of Python—2 or 3—on the same machine using “environments” with the `conda` tool described in Section A.1.2.

APPENDIX F

Under the Hood

This appendix describes how Python handles variables and objects internally. It is not essential to understand this material to complete the exercises in this tutorial, but it may be useful when analyzing errors and writing (or reading) more advanced code in the future.

F.1 ASSIGNMENT STATEMENTS

Section 2.1 claimed that in Python, everything is an object, endowed with data fields and methods. When executing an assignment like `x=np.arange(10)`, Python binds a variable name to an object.

In some programming languages, a statement corresponding to `x=np.arange(10)` would allocate a block of memory, assign it the name `x`, then place an array of integers inside that block. The block of memory is permanently linked to the name `x`, so the name and the object it represents are essentially the same thing. This is not true in Python. Python creates an `ndarray` object, then stores its memory address in `x`. Thus, the variable `x` *points to* the `ndarray` object. The name `x` can be used to access all of the methods and data fields of the `ndarray` object, but the `ndarray` object and the variable `x` are separate entities.

One consequence of this arrangement is that two variables can point to the same object (the same memory address). Try this:

```
x = np.zeros(10)
y = x
x[1] = 1
y[0] = 1
5 print("x={}\ny={}".format(x, y))
```

You'll see that `x` and `y` both point to the same array. Assignment statements can also bind an existing variable to a different object. Try the following lines of code:

```
x = np.zeros(10)
y = x
y = y + 1
```

After the first two lines, `x` and `y` point to the same array. The assignment in the third line creates a new array to hold the result of `y+1` and then binds the variable `y` to the new array. It has no effect on `x` or the original array. (You can verify this with a couple of `print` commands.)

An assignment statement like `y=x` does not permanently link `x` and `y`. They only point to the same object until one of them is reassigned.

A consequence of this rule is that an assignment statement like `y=x+1` does not link `x` and `y` at all: `y` points to a new object created after evaluating the expression `x+1`.

You can use built-in Python functions to determine whether variables point to the same object. The `id` function will return the address in your computer's memory that a variable points to. If `id(x)` and `id(y)` are different, then `x` and `y` point to different objects. Likewise, `x is y` will only return `True` if `x` and `y` refer to the same object. Try the following commands:

```
x = np.arange(10)
y = x
z = np.arange(10)

5 print("x==y: ", x==y)
print("x==z: ", x==z)
print("x is y: ", x is y)
print("x is z: ", x is z)

10 print("id(x): ", id(x))
print("id(y): ", id(y))
print("id(z): ", id(z))
```

Even though all of the elements of `x`, `y`, and `z` are the same, we see that `x` and `y` are actually the same array object, while `z` is a separate array object that happens to have the same entries.

If `y=x` binds both objects to the same memory address, how can you make an independent copy of an object? The answer depends on the type of object. You can copy a Python list by slicing:

```
x = [1, 1, 1]
y = x[:]
x == y
```

The variables `x` and `y` now contain the same elements, but are they the same list? Find out:

```
x[1] = 0
print("x={}\ny={}".format(x, y))
```

You *cannot* copy a NumPy array by slicing, however. Slicing creates a new `view` of the existing array data, not a copy.¹ Thus, modifying an element of the array also modifies any slice containing that element. Likewise, modifying a slice will change the original array. To create an independent copy of an array, you must use a method called `copy`. To understand the difference between a slice and a copy, try this:

```
w = np.zeros(10)
x = w
y = x[:]
z = y.copy()
5 y[0] = 1                      # Modify a slice.
z[1] = 1                      # Modify a copy.
print("w={}\nx={}\ny={}\nz={}".format(w, x, y, z))
```

The relation between variables and objects and the effects of assignment statements are illustrated in Figure F.1.

¹ The array methods `ravel` and `reshape` mentioned in Section 2.2.9 also return a new view of an existing array.

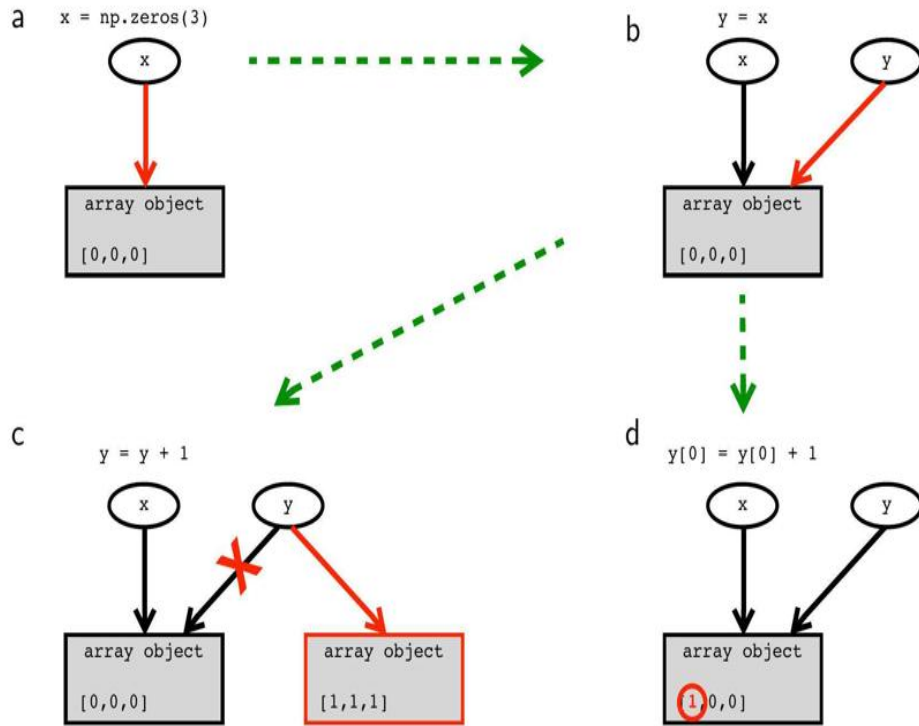


Figure F.1: Relationship between variables and objects in Python. The effects of the assignment statements are shown in red. (a) *Assigning a variable to a new object*. The call to `np.zeros(3)` creates an array object. Python then creates a variable named `x` that points to the array. (b) *Assigning a variable to an existing object*. The assignment statement `y=x` does not create a copy of `x`. Instead, Python binds the variable `y` to the *same* array object as `x`. (c) *Reassignment of a variable*. After (b), both variables point to the same array. In the assignment `y=y+1`, Python creates a new array when it evaluates `y+1`. It then binds `y` to this new array, with no effect on `x`. (d) *Changing an element of an array*. In contrast to (c), the statement `y[0]=y[0]+1` changes the value stored at position 0 of the array that `y` points to. Because `x` and `y` still point to the same object, `x[0]` also changes.

F.2 MEMORY MANAGEMENT

The objects created by Python do not persist forever. Python has a *garbage collection* routine that periodically checks whether *any* variables point to each object for which it has allocated memory. If no variables are bound to an object, then that object is deleted and its memory is reclaimed.

You can forcibly remove a *variable* by using the `del` statement: `del(x)`. However, Python will not destroy the object `x` was pointing to unless *no* variables point to that object. Memory management is under Python's control, not yours. In general, this is a good thing.

F.3 FUNCTIONS

When a variable is passed to a function, what exactly is passed?

In computer science, there are two common paradigms: *pass by value* and *pass by reference*. Pass by value means a function receives a copy of an object passed as an argument. Because the function acts on a copy, *nothing* it does to its argument affects the original object. In contrast, pass by reference means a function receives the memory address of an object passed as an argument. Because the function acts on the original object, *everything* it does to its argument affects the original object. If f is pass by value, then $f(x)$ has no effect on x . If f is pass by reference, then $f(x)$ can affect x .

Consider the following example:

```

def object_plus_one(y):
    y = y + 1

def elements_plus_one(y):
5     for i in range(len(y)):
        y[i] = y[i] + 1

x = np.ones(10)
object_plus_one(x)
10 print(x)
elements_plus_one(x)
print(x)

```

One function changes x ; the other does not. It appears that Python uses both paradigms, even though we did not specify either when writing the functions! How can you predict what will happen when you write a function? The key is to remember that a variable *points to* an object, but it is *not* the object. Technically, arguments to functions are always passed by value in Python. However, the “value” being passed is a reference to an object (an address in memory). Thus, a function receives a *copy* of the memory address where an argument’s object is stored. It creates a new local variable for each argument and binds it to the same object as the corresponding argument. A function $f(x)$ cannot modify this address and make $f(x)$ point to a *different* object, but it can use this address to access methods of the object stored there, and thereby modify the object that x points to. To distinguish this behavior from the ordinary usage of pass by value and pass by reference, the paradigm used in Python is often referred to as *reference passed by value*, or *pass by address*.

To better understand the behavior of functions, we next need to explore how Python finds an object when it encounters a variable name.

F.4 SCOPE

The way in which Python looks up the value of a variable is subtle but important. Python keeps track of variables using namespaces. A **namespace** is like a directory of names and objects that tells Python where to find the object associated with a variable name. The subtle point is that Python maintains multiple namespaces, each with its own **scope**. A scope is a portion of a program, like a function body or a module, in which a namespace is accessible. For example, there is a namespace that keeps track of all the variables you define at the command prompt. Its scope is *global*: Any script you run or commands you type at the command prompt can find and use these variables.

Each time Python executes a function, it creates a *local* namespace containing all variables created

within the function. They are concealed from everything outside, because the scope of the local namespace

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is limited to the function itself. Recall the `taxicab` function of the `measurements.py` module in Section 6.1:

```
# excerpt from measurements.py      [get code]
def taxicab(pointA, pointB):
    """
    Taxicab metric for computing distance between points A and B.
    pointA = (x1, y1)
    pointB = (x2, y2)
    Returns |x2-x1| + |y2-y1|. Distances are measured in city blocks.
    """
    interval = abs(pointB[0] - pointA[0]) + abs(pointB[1] - pointA[1])
    return interval
10
```

There is no variable called `interval` in the Variable Explorer before or after you execute the `taxicab` function. It only exists in the local namespace of the function.

When you refer to a variable from within a function, Python will look for an object with that name in an expanding search. Suppose that you accidentally typed `x1` instead of `pointA[0]` in the body of `taxicab`. When you call the function, Python will search for the name `x1` in the following namespaces, in order:

Local: First, Python determines whether `x1` is defined in the function body or in its argument list. If so, it ends the search and uses the object bound to that variable. There is no variable called `x1` in this example, so Python expands the search.

Enclosing: Next, Python will determine whether the current function is defined within another function. If so, it will determine whether `x1` was defined within or passed as an argument to this or any other *enclosing* functions. There may be multiple nested functions, each with its own namespace for Python to search.

Global: If it still has not found `x1`, Python will check to see whether the module in which the function is defined contains a variable called `x1`. (For example, if you had added the line `x1=10` outside of any function definitions in the `measurements.py` module, Python would stop searching and use this value. If you defined `taxicab` at the command prompt or by running a script instead of importing the module, then Python would search among all the variables in your current session.)

Built-in: As a last resort, Python will check its built-in functions and parameters—those listed in `dir(__builtins__)`. If it cannot find `x1` here, it gives up and raises a `NameError`.

Knowing this hierarchy of $L \rightarrow E \rightarrow G \rightarrow B$ can be useful in debugging. Python may be finding a value for a variable somewhere you never intended.

The following function demonstrates Python's rules of scope.

```
# scope.py      [get code]
def illustrate_scope():
    s_enclosing = 'E'
    def display():
        s_local = 'L'
5
```

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```

    print("Local --> {}".format(s_local))
    print("Enclosing --> {}".format(s_enclosing))
    print("Global --> {}".format(s_global))
10   print("Built-in --> {}".format(abs))

display()

s_global = 'G'
illustrate_scope()

```

This script defines three variables whose names indicate their namespaces with respect to the `display()` function. When `illustrate_scope()` is called in the final line, Python executes the function body. It defines a variable in the current namespace, defines the `display()` function, and then calls this newly defined function. When Python executes `display()`, it must search for four different names. It finds `s_local` within the local namespace, but none of the other variables are defined there. Python finds `s_enclosing` in the enclosing namespace, `s_global` in the global namespace, and `abs` within Python's collection of built-in functions.

F.4.1 Name collisions

Scope is important in determining the effect of a function call. In the preceding example, all of the variables had unique names. But what happens when variables in different namespaces have the same name? Try the following example to see how Python resolves name collisions.

```

# name_collision.py      [get code]
def name_collisions():
    x, y = 'E', 'E'
    def display():
        x = 'L'
        print("Inside display() ...")
        print("x= {}\ny= {}\nz= {}".format(x, y, z))
    display()
    print("Inside name_collision() ...")
10   print("x= {}\ny= {}\nz= {}".format(x, y, z))

    x, y, z = 'G', 'G', 'G'
    name_collisions()
    print("Outside function ...")
15   print("x= {}\ny= {}\nz= {}".format(x, y, z))

```

Each assignment statement creates a variable in the current namespace, but it has no effect on the variables in other namespaces. Thus, in this example there are three variables with the name `x`, two with the name `y`, and one with the name `z`. When Python executes `name_collision()` and `display()`, it must determine a value for each variable. It starts searching in the innermost namespace, works its way through the $L \rightarrow E \rightarrow G \rightarrow B$ hierarchy, and uses the value from the first namespace in which it locates each variable name.

Python uses namespaces, each with its own scope, to protect programs from unintended name collisions. This means that you do not have to worry about giving each variable a globally unique name when writing your own functions. However, Python cannot help you if you use the same variable name

for two different purposes in the same namespace, as discussed in Section 1.4.3. Reassigning a name destroys any previous connection to another object. (Python’s namespaces do not include the mind of the programmer!) Modular programming—breaking complex programs into a series of simple functions that each accomplish one thing—and descriptive variable names are the best ways to prevent this type of name collision.

F.4.2 Variables passed as arguments

As mentioned in Section F.3, when a variable is passed to a function as an argument, Python creates a new variable in the function’s local namespace and binds it to the same object as the argument. Despite referring to the same object, these two variables are independent—even if they have the *same name!* They exist in different namespaces.

If an assignment statement within a function binds an argument variable to a new value, there is no effect on the global or enclosing variable passed as that argument. Python simply reassigns the local variable to a different object. In this way, Python allows functions to *access* external variables, not *modify* them. Such modularity enforces good coding practice; however, there is an important exception to this principle. A function *can* modify an object by using that object’s methods. The more precise rule is, “A function cannot *reassign* an external variable.” If a function modifies an argument using a method of that argument, it is modifying an object that is bound to variables in multiple namespaces. This will result in side effects—whether by design or mistake.

We can now explain the behavior of `object_plus_one(x)` and `elements_plus_one(x)` in Section F.3. When `x` is passed as an argument to the function, a new local variable `y` is created in the function’s local namespace and bound to the same array as `x`, as in Figure F.1b. The assignment `y=y+1` binds the local `y` to a new object with no effect on the global `x`, as in Figure F.1c. Because no value is returned by the function, and Python deletes local variables after evaluating the function, `object_plus_one(x)` has no external effects—it accomplishes nothing.

In contrast, `elements_plus_one(x)` does not reassign its local variable `y` to a new object. It uses an array method to modify the data of `y`. (See Section 2.2.6, page 26.) The statement `y[0]=y[0]+1` changes the value of the first element of the array, as shown in Figure F.1d. Because the variable `y` in the function’s local namespace and the variable `x` in the global namespace point to the *same* array object, `x[0]` is also changed. This continues as the function iterates over the loop. Unlike `object_plus_one(x)`, `elements_plus_one(x)` produces a side effect: It changes the value of `x`.

F.5 SUMMARY

When we think about an assignment statement like `x=2.0`, it’s natural to imagine that some memory is set aside for `x`, then the value 2.0 is stored there. Python does not use this approach. Instead, it creates a `float` object with the value 2.0, and then binds the variable `x` to that object.

In Python, objects and variables exist independently of one another. After an assignment statement, a variable points to the address in memory where an object is stored.

Variables have a scope: They can only be accessed by certain portions of a program, and Python uses

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namespaces to prevent conflicts between variables with the same name. There is no such protection for the objects they are bound to. Multiple variables can point to the same object, and this can lead to unexpected results when you are trying to carry out different operations on what you thought were copies of the same data. Functions create temporary variables that are initially bound to the same objects as their arguments. This, too, can lead to unexpected behavior.

Errors due to improper use of Python's system of variables bound to objects can be difficult to diagnose. The best way to avoid them is to plan carefully when writing code. When you make an assignment or write a function, consider: Do I want two variables that point to the *same* data? Or do I want to make a *copy* of the data?

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APPENDIX G

Answers to “Your Turn” Questions

Your Turn 2B (page 23):

The `np.array` function is called on a list of lists—a two-element list whose elements are themselves three-element lists of integers—so it creates a 2×3 array to store the data. Python starts filling in the $(0, 0)$ cell of the array. Each comma moves to the next column in the same row of the array. At the end of the first list, that is, after “`J`”, Python moves to the first column of the next row of the array. The resulting array is equivalent to a matrix with two rows and three columns containing the original data.

Your Turn 2C (page 24):

The `np.linspace` function ends exactly at 10 and uses whatever spacing it needs to accomplish that. In contrast, `np.arange` uses the exact spacing of 1.5 and ends just before it reaches 10.

Your Turn 2D (page 27):

To get the odd-index elements, use `a[1::2]`. This instructs Python to start at offset 1, then step by 2 until it reaches the end of the array.

Your Turn 2E (page 30):

Python concatenates three strings to produce the output: `s`, a single space, and `t`.

Your Turn 3A (page 39):

a.

```
x = np.linspace(-3, 3, 101)
y = np.exp(-x**2)
```

b.

```
from scipy.special import factorial
mu, N = 2, 10
n_array = np.arange(N + 1)
poisson = np.exp(-mu) * (mu**n_array) / factorial(n_array)
```

Your Turn 3B (page 40):

`a*b` attempts item-by-item evaluation, but the shapes don’t match. However, because `a` has only one row, Python uses NumPy’s broadcasting rules and uses that row as many times as needed. Similarly, because `b` has only one column, Python uses that column as many times as needed. The result is therefore a 3×3 array. `np.dot(a, b)` follows the usual matrix rules, arriving at a 2D array with just one row and one column.

Your Turn 4A (page 63):

On a semilog plot, the exponential function appears as a straight line. Power laws do not.

On a log-log plot, power laws appear as straight lines; exponentials do not.

```
x = np.linspace(2, 7, 51)
nu = 3.6

plt.figure()
5 plt.plot(x, np.exp(x), label='Exponential')
plt.plot(x, x**nu, label='Power Law')
plt.title("Linear Plot")
plt.legend()

10 plt.figure()
plt.semilogy(x, np.exp(x), label='Exponential')
plt.semilogy(x, x**nu, label='Power Law')
plt.title("Semilog Plot")
plt.legend()

15 plt.figure()
plt.loglog(x, np.exp(x), label='Exponential')
plt.loglog(x, x**nu, label='Power Law')
plt.title("Log-Log Plot")
20 plt.legend()
```

Your Turn 4B (page 65):

```
# fancy_plot.py      [get code]
x_min, x_max = 0, 4
num_points = 51
x_vals = np.linspace(x_min, x_max, num_points)
5 y_vals = x_vals**2
plt.plot(x_vals, y_vals, 'r', linewidth=3)

ax = plt.gca()
ax.set_title("My Little Plot", fontsize=32)
10 ax.set_xlabel("$x$", fontsize=24)
ax.set_ylabel("$y = x^2$", fontsize=24)
```

Note that we have to set the font size for each element of the figure separately. (Chapter 10 gives an elegant alternative.)

Your Turn 4C (page 67):

To add a legend to the existing plot, use

```
# legend.py      [get code]
ax = plt.gca()
ax.legend( ("sin(2$\backslash\pi$x)", \
5           "sin(4$\backslash\pi$x)", \
           "sin(6$\backslash\pi$x)" )
```


The rest of the embellishment is up to you!

Your Turn 6A (page 77):

The function should be similar to `taxicab` with a modified distance function. The docstring should specify that each point contain three elements.

```
# measurements.py      [get code]
def crow(pointA, pointB):
    """
    Distance between points A and B "as the crow flies."
    5      pointA = (x1, y1, z1)
           pointB = (x2, y2, z2)
    Returns sqrt((x2-x1)**2 + (y2-y1)**2 + (z2-z1)**2)
    """
    10     distance = np.sqrt( (pointA[0] - pointB[0])**2 + \
                           (pointA[1] - pointB[1])**2 + \
                           (pointA[2] - pointB[2])**2 )
    return distance
```

Your Turn 6B (page 83):

- a. `x_step = 2*x_step - 1`
- b. `x_position = np.cumsum(x_step)` or `x_position = x_step.cumsum()`
- c.

```
# random_walk.py      [get code]
# Create a random number generator.
rng = np.random.default_rng() # create a random number generator object
rand = rng.random           # assign its uniform distribution method to rand
5
num_steps = 1000

#%% Plot a single random walk.
x_step = rand(num_steps) > 0.5
10 y_step = rand(num_steps) > 0.5
x_step = 2*x_step - 1
y_step = 2*y_step - 1
x_position = np.cumsum(x_step)
y_position = np.cumsum(y_step)
15 plt.figure()
plt.plot(x_position, y_position)
plt.axis('square')

20 %% Plot a grid of random walks.
M, N = 4, 4
fig, ax = plt.subplots(M, N, sharex=True, sharey=True)

for i in range(4):
```



```

25     for j in range(4):
26         x_step = rand(num_steps) > 0.5
27         y_step = rand(num_steps) > 0.5
28         x_step = 2*x_step - 1
29         y_step = 2*y_step - 1
30         x_position = np.cumsum(x_step)
31         y_position = np.cumsum(y_step)
32         ax[i,j].plot(x_position, y_position)

```

Your Turn 6C (page 84):

By default, `plt.hist` and `np.histogram` set up ten equally spaced bins. The variable `bin_edges` is an array with *eleven* elements—the *edges* of the bins. The first element is the smallest entry in the data list; the last element is the largest entry in the data list. Each bin's width is `(data.max() - data.min()) / 10`. Meanwhile, `counts[i]` contains all the elements of `data` that fall between `bin_edges[i]` and `bin_edges[i+1]`. Thus, `bin_edges` will always contain one more element than `counts`.

Your Turn 6D (page 87):

```

# surface.py      [get code]
from mpl_toolkits.mplot3d import Axes3D
points = np.linspace(-1, 1, 101)
X, Y = np.meshgrid(points, points)
Z = X**2 + Y**2
ax = Axes3D(plt.figure())
ax.plot_surface(X, Y, Z)
# ax.plot_surface(X, Y, Z, rcount=10, ccount=10)
# ax.plot_surface(X, Y, Z, rstride=1, cstride=1)

```

This script uses the default mesh. The commented lines give different meshes. The second to last line produces a coarse surface; the final line produces a smooth surface by using every data point.

Your Turn 6F (page 93):

a.

```

from scipy.integrate import quad
def f(x): return x**2
integral, error = quad(f, 0, 2)
print("Difference from exact result: ", integral - 2**3 / 3)

```

b.

```

from scipy.integrate import quad
x_max = np.linspace(0, 5, 51)
integral = np.zeros(x_max.size)
def integrand(x): return np.exp(-x**2/2)
for i in range(x_max.size):
    integral[i], error = quad(integrand, 0, x_max[i])
plt.plot(x_max, integral)

```


C.

```
from scipy.integrate import quad
def integrand(x): return np.exp(-x**2/2)
integral, error = quad(integrand, -np.inf, np.inf)
print("Difference from exact result: ", integral - np.sqrt(2*np.pi))
```

Your Turn 6G (page 97):

Driving the system near its resonant frequency ($\omega_0 = 1$) yields a strong response. The driven mode of frequency $\omega = 0.8$ is superposed with the natural modes required to satisfy the initial conditions. Interference between modes of different frequencies produces a pattern of beats.

Your Turn 6H (page 100):

The vector at the point (x, y) has components $(y, -x)$. This arrow is always perpendicular to the vector from the origin to its base point, just like the velocity vector field of a rigid, spinning disk.

Your Turn 6I (page 102):

- This example gives trajectories that all spiral into the origin, because we added a small component to \mathbf{V} that points radially inward.
- This example gives a fixed point at the origin of “saddle” type. One of the initial conditions runs smack into the fixed point at the origin, but the others veer away and run off to infinity.

Your Turn 8A (page 110):

The resulting image is a negative of the original with no shades of gray, just black and white. Everything darker than the mean lightness became fully white, and everything lighter than the mean became fully black. In performing the array operation, NumPy has modified the data type of the array from `uint8` to `bool`. PyPlot has no trouble generating images and saving files with arrays of different data types, but the array is no longer a collection of 8-bit luminance values.

Your Turn 9A (page 119):

- In the Python indexing scheme, the only allowed index for F is $(k, \ell) = (0, 0)$. Thus, $C_{i,j} = F_{0,0} \cdot I_{i,j} = I_{i,j}$.
- Again using the Python indexing scheme, the allowed indices in Equation 9.1 run from $(i, j) = (0, 0)$ with $(k, \ell) = (0, 0)$ to $(i, j) = (M + m - 2, N + n - 2) = ([M - 1] + [m - 1], [N - 1] + [n - 1])$ with $(k, \ell) = (m - 1, n - 1)$. Thus, C contains $(M + m - 1) \times (N + n - 1)$ entries.

Your Turn 10A (page 130):

Follow the pattern of `data_dictionary.py` using different input parameters.

Your Turn 10B (page 135):

The following code reveals the problem.

```
N = 10**4
samples = 10**4
targets = [10, 20, 50, 100, 200, 500]

5 df = pd.DataFrame()
```



```

for L in targets:
    df["L={}".format(L)] = [first_passage(L,N) for n in range(samples)]

df.hist(sharey=True)
10 df.plot(kind='density', legend=True)
plt.legend()

```

As the target distance increases, the number of walks that do not reach the target distance increases rapidly. The changing shapes of the histograms and probability densities reveal that the tail of the distribution is being cut off. Fewer valid samples imply noisier statistics, and the average of the samples that did reach the target is no longer representative of the probability distribution we sought to study.

Repeating the simulations with $N=10^{*6}$ and $\text{samples}=10^{*5}$ improves the statistics—especially for longer target distances.

Your Turn 10C (page 138):

The data follow a clear linear relationship. Even the power law fit gives an exponent of 1. One must be careful to make the number of steps in the walk large enough or the target distance small enough, though. We ignore walks that do not reach the target, but this will bias the average downward and reduce the number of data points at large values of L . Even 1 million steps is insufficient for about 40 percent of the walkers to reach $L = 500$.

Because of the large number of steps and walks, this script takes a while to run.

```

# regression.py      [get code]
import numpy as np
import matplotlib.pyplot as plt
import pandas as pd
5 from sklearn.linear_model import LinearRegression

# Define and run simulations.
data = {}          # empty dictionary to store all data
nmax = 10**6
10 parameters = dict(N=nmax, p=0.5)  # common inputs
l_values = [10, 20, 50, 100, 200, 500]
samples = 10000

for l in l_values:
15     data["L={}".format(l)] = \
        [first_passage(L=l, **parameters) for n in range(samples)]
step_data = pd.DataFrame(data)

# Prepare data for modeling with sklearn.
20 model = LinearRegression()

X = np.array(l_values).reshape(-1,1)
Y = step_data.mean()
logX = np.log(X)
25 logY = np.log(Y)
xFit = np.linspace(1, X.max(), 201).reshape(-1,1)

```

```
# Plot data.
plt.figure()
30 plt.plot(X, Y, 'ko', mew=2, mfc='none', label="Data")

# Check for linear relationship.
model.fit(X, Y)
print("Linear Model: A + b*x")
35 print("A = {:.3f}, b = {:.3f}".format(model.intercept_, model.coef_[0]))
print("R2: ", model.score(X,Y))
yFit = model.predict(xFit)
plt.plot(xFit, yFit, 'r-', label="Linear Model")

40 # Check for exponential relationship.
model.fit(X, logY)
print("Exponential Model: A * exp(b*x)")
print("A = {:.3f}, b = {:.3f}".format(model.intercept_, model.coef_[0]))
print("R2: ", model.score(X,Y))
45 yFit = model.predict(xFit)
plt.plot(xFit, np.exp(yFit), 'g-', label="Exponential Model")

# Check for power-law relationship.
model.fit(logX, logY)
50 print("Power Law Model: A * x**b")
print("A = {:.3f}, b = {:.3f}".format(model.intercept_, model.coef_[0]))
print("R2: ", model.score(X,Y))
yFit = model.predict(np.log(xFit))
plt.plot(xFit, np.exp(yFit), 'b-', label="Power Law Model")
55
plt.legend()
```

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Acknowledgments

Many students and colleagues taught us tricks that ended up in this tutorial, especially Tom Dodson. Alexander Alemi, Steve Baylor, Gary Bernstein, Kevin Chen, R. Michael Jarvis, Michael Lerner, Dave Pine, and Jim Sethna gave expert advice and caught many errors. Three anonymous reviewers not only gave us expert suggestions, but also questioned sharply the goals of the book in ways that helped us to clarify those goals.

Nily Dan reminded us to strive for utility, not erudition. Kim Kinder provided encouragement and diversion in equal measure, at just the right times, throughout the writing of this book.

This tutorial was set in \LaTeX using the `listings` package. Jobst Hoffman and Heiko Oberdiek kindly supplied personal help. (The package settings for our code listings are available at the book's blog.)

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Recommended Reading

If you can know everything about anything, it is not worth knowing.
— Robertson Davies

Starting from the foundation in this tutorial, you may be able to get the advanced material you need for a particular problem just from web searches. Some other references that we found helpful appear below.

As your skills develop, you may start writing longer and more complex codes. Then it will pay off to make an additional investment in learning about basic software engineering practices (for example, in Scopatz & Huff, 2015) and specifically about user-defined classes (for example, in Guttag, 2021; Scopatz & Huff, 2015). The PEP 8 Style guide for Python also provides guidelines for writing Python code in a standard, readable way: www.python.org/dev/peps/pep-0008/.

This book has alluded to the free L^AT_EX typesetting system, in the context of graph labels and Jupyter notebooks. You can obtain it, and its documentation, from www.latex-project.org/get/.

A blog accompanies this book. It can be accessed via press.princeton.edu/titles/32489.html, or directly at physicalmodelingwithpython.blogspot.com/. Here you will find data sets, code samples, errata, additional resources, and extended discussions of the topics introduced in this book.

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