Practical Programming for Biological Data

Covers Pation 3



Using Python



Powerful, flexible, and easy to use, Python is an ideal language for building software tools and applications for life science research and development. This unique book shows you how to program with Python, using code examples taken directly from bioinformatics. In a short time, you'll be using sophisticated techniques and Python modules that are particularly effective for bioinformatics programming.

Bioinformatics Programming Using Python is perfect for anyone involved with bioinformatics—researchers, support staff, students, and software developers interested in writing bioinformatics applications. You'll find it useful whether you already use Python, write code in another language, or have no programming experience at all. It's an excellent self-instruction tool, as well as a handy reference when facing the challenges of real-life programming tasks.

- Become familiar with Python's fundamentals, including ways to develop simple applications
- Learn how to use Python modules for pattern matching, structured text processing, online data retrieval, and database access
- Discover generalized patterns that cover a large proportion of how Python code is used in bioinformatics
- Learn how to apply the principles and techniques of object-oriented programming
- Benefit from the "tips and traps" section in each chapter

"Mitchell Model has done the bioinformatics community a real service. In biology, already an information-heavy science, new generations of sequencers have opened the floodgates of data. The Python programming techniques introduced here will enable you to transform that data into knowledge."

—Jim Tisdall

Author of Beginning Perl
for Bioinformatics

Mitchell L Model has been focused on bioinformatics for 15 years and has worked with a wide range of platforms, languages, technologies, and domains. As an independent consultant, he provides training, mentoring, tools, and support to software development groups learning new technologies and practices.



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Mitchell L Model



by Mitchell L Model

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Table of Contents

Prefa	ce	. xi
1.	Primitives	. 1
	Simple Values	1
	Booleans	2
	Integers	2 2 3
	Floats	3
	Strings	4
	Expressions	5
	Numeric Operators	5 7
	Logical Operations	
	String Operations	9
	Calls	12
	Compound Expressions	16
	Tips, Traps, and Tracebacks	18
	Tips	18
	Traps	20
	Tracebacks	20
2.	Names, Functions, and Modules	21
	Assigning Names	23
	Defining Functions	24
	Function Parameters	27
	Comments and Documentation	28
	Assertions	30
	Default Parameter Values	32
	Using Modules	34
	Importing	34
	Python Files	38
	Tips, Traps, and Tracebacks	40
	Tips	40

	Traps	45
	Tracebacks	46
3.	Collections	47
	Sets	48
	Sequences	51
	Strings, Bytes, and Bytearrays	53
	Ranges	60
	Tuples	61
	Lists	62
	Mappings	66
	Dictionaries	67
	Streams	72
	Files	73
	Generators	78
	Collection-Related Expression Features	79
	Comprehensions	79
	Functional Parameters	89
	Tips, Traps, and Tracebacks	94
	Tips	94
	Traps	96
	Tracebacks	97
4.	Control Statements	99
	Conditionals	101
	Loops	104
	Simple Loop Examples	105
	Initialization of Loop Values	106
	Looping Forever	107
	Loops with Guard Conditions	109
	Iterations	111
	Iteration Statements	111
	Kinds of Iterations	113
	Exception Handlers	134
	Python Errors	136
	Exception Handling Statements	138
	Raising Exceptions	141
	Extended Examples	143
	Extracting Information from an HTML File	143
	The Grand Unified Bioinformatics File Parser	146
	Parsing GenBank Files	148
	Translating RNA Sequences	151
	Constructing a Table from a Text File	155

	Tips, Traps, and Tracebacks	160
	Tips	160
	Traps	162
	Tracebacks	163
5.	Classes	. 165
	Defining Classes	166
	Instance Attributes	168
	Class Attributes	179
	Class and Method Relationships	186
	Decomposition	186
	Inheritance	194
	Tips, Traps, and Tracebacks	205
	Tips	205
	Traps	207
	Tracebacks	208
6.	Utilities	. 209
	System Environment	209
	Dates and Times: datetime	209
	System Information	212
	Command-Line Utilities	217
	Communications	223
	The Filesystem	226
	Operating System Interface: os	226
	Manipulating Paths: os.path	229
	Filename Expansion: fnmatch and glob	232
	Shell Utilities: shutil	234
	Comparing Files and Directories	235
	Working with Text	238
	Formatting Blocks of Text: textwrap	238
	String Utilities: string	240
	Comma- and Tab-Separated Formats: csv	241
	String-Based Reading and Writing: io	242
	Persistent Storage	243
	Persistent Text: dbm	243
	Persistent Objects: pickle	247
	Keyed Persistent Object Storage: shelve	248
	Debugging Tools	249
	Tips, Traps, and Tracebacks	253
	Tips	253
	Traps	254
	Tracebacks	2.5.5

7.	Pattern Matching	. 257
	Fundamental Syntax	258
	Fixed-Length Matching	259
	Variable-Length Matching	262
	Greedy Versus Nongreedy Matching	263
	Grouping and Disjunction	264
	The Actions of the re Module	265
	Functions	265
	Flags	266
	Methods	268
	Results of re Functions and Methods	269
	Match Object Fields	269
	Match Object Methods	269
	Putting It All Together: Examples	270
	Some Quick Examples	270
	Extracting Descriptions from Sequence Files	272
	Extracting Entries From Sequence Files	274
	Tips, Traps, and Tracebacks	283
	Tips	283
	Traps	284
	Tracebacks	285
8.	Structured Text	. 287
	HTML	287
	Simple HTML Processing	289
	Structured HTML Processing	297
	XML	300
	The Nature of XML	300
	An XML File for a Complete Genome	302
	The ElementTree Module	303
	Event-Based Processing	310
	expat	317
	Tips, Traps, and Tracebacks	322
	Tips	322
	Traps	323
	Tracebacks	323
9.	Web Programming	. 325
	Manipulating URLs: urllib.parse	325
	Disassembling URLs	326
	Assembling URLs	327
	Assembling ORLs	321
		328
	Opening Web Pages: webbrowser Module Functions	

	Constructing and Submitting Queries	329
	Constructing and Viewing an HTML Page	330
	Web Clients	331
	Making the URLs in a Response Absolute	332
	Constructing an HTML Page of Extracted Links	333
	Downloading a Web Page's Linked Files	334
	Web Servers	337
	Sockets and Servers	337
	CGI	343
	Simple Web Applications	348
	Tips, Traps, and Tracebacks	354
	Tips	355
	Traps	357
	Tracebacks	358
10.	Relational Databases	. 359
	Representation in Relational Databases	360
	Database Tables	360
	A Restriction Enzyme Database	365
	Using Relational Data	370
	SQL Basics	371
	SQL Queries	380
	Querying the Database from a Web Page	392
	Tips, Traps, and Tracebacks	395
	Tips	395
	Traps	398
	Tracebacks	398
11.	Structured Graphics	. 399
	Introduction to Graphics Programming	399
	Concepts	400
	GUI Toolkits	404
	Structured Graphics with tkinter	406
	tkinter Fundamentals	406
	Examples	411
	Structured Graphics with SVG	431
	SVG File Contents	432
	Examples	436
	Tips, Traps, and Tracebacks	444
	Tips	444
	Traps	445
	Tracebacks	447

A.	Python Language Summary	449
В.	Collection Type Summary	459
Index		473

Preface

This preface provides information I expect will be important for someone reading and using this book. The first part introduces the book itself. The second talks about Python. The third part contains other notes of various kinds.

Introduction

I would like to begin with some comments about this book, the field of bioinformatics, and the kinds of people I think will find it useful.

About This Book

The purpose of this book is to show the reader how to use the Python programming language to facilitate and automate the wide variety of data manipulation tasks encountered in life science research and development. It is designed to be accessible to readers with a range of interests and backgrounds, both scientific and technical. It emphasizes practical programming, using meaningful examples of useful code. In addition to meeting the needs of individual readers, it can also be used as a textbook for a one-semester upper-level undergraduate or graduate-level course.

The book differs from traditional introductory programming texts in a variety of ways. It does not attempt to detail every possible variation of the mechanisms it describes, emphasizing instead the most frequently used. It offers an introduction to Python programming that is more rapid and in some ways more superficial than what would be found in a text devoted solely to Python or introductory programming. At the same time, it includes some advanced features, techniques, and topics that are often omitted from entry-level Python books. These are included because of their wide applicability in bioinformatics programming, and they are used extensively in the book's examples.

Python's installation includes a large selection of optional components called "modules." Python books usually cover a small selection of the most generally useful modules, and perhaps some others in less detail. Having bioinformatics programming as this book's target had some interesting effects on the choice of which modules to discuss, and at what depth. The modules (or parts of modules) that are

covered in this book are the ones that are most likely to be particularly valuable in bioinformatics programming. In some cases the discussions are more substantial than would be found in a generic Python book, and many of the modules covered here appear in few other books. Chapter 6, in particular, describes a large number of narrowly focused "utility" modules.

The remaining chapters focus on particular areas of programming technology: pattern matching, processing structured text (HTML and XML), web programming (opening web pages, programming HTTP requests, interacting with web servers, etc.), relational databases (SQL), and structured graphics (Tk and SVG). They each introduce one or two modules that are essential for working with these technologies, but the chapters have a much larger scope than simply describing those modules.

Unlike many technical books, this one really should be read linearly. Even in the later chapters, which deal extensively with particular kinds of programming work, examples will often use material from an earlier chapter. In most places the text says that and provides cross-references to earlier examples, so you'll at least know when you've encountered something that depends on earlier material. If you do jump from one place to another, these will provide a path back to what you've missed.

Each chapter ends with a special "Tips, Traps, and Tracebacks" section. The tips provide guidance for applying the concepts, mechanisms, and techniques discussed in the chapter. In earlier chapters, many of the tips also provide advice and recommendations for learning Python, using development tools, and organizing programs. The traps are details, warnings, and clarifications regarding common sources of confusion or error for Python programmers (especially new ones). You'll soon learn what a traceback is; for now it is enough to say that they are error messages likely to be encountered when writing code based on the chapter's material.

About Bioinformatics

Any title with the word "bioinformatics" in it is intrinsically ambiguous. There are (at least) three quite different kinds of activities that fall within this term's wide scope. Both the nature of the work performed and the educational backgrounds and technical talents of the people who perform these various activities differ significantly. The three main areas of bioinformatics are:

Computational biology

Concerned with the development of algorithms for mining biological data and modeling biological phenomena

Software development

Focused on writing software to implement computational biology algorithms, visualize complex data, and support research and development activity, with particular attention to the challenges of organizing, searching, and manipulating enormous quantities of biological data

Life science research and development

Focused on the application of the tools and results provided by the other two areas to probe the processes of life

This book is designed to teach you bioinformatics software development. There is no computational biology here: no statistics, formulas, equations—not even explanations of the algorithms that underlie commonly used informatics software. The book's examples are all based on the kind of data life science researchers work with and what they do with it.

The book focuses on practical data management and manipulation tasks. The term "data" has a wide scope here, including not only the contents of databases but also the contents of text files, web pages, and other information sources. Examples focus on genomics, an area that, relative to others, is more mature and easier to introduce to people new to the scientific content of bioinformatics, as well as dealing with data that is more amenable to representation and manipulation in software. Also, and not incidentally, it is the part of bioinformatics with which the author is most familiar.

About the Reader

This book assumes no prior programming experience. Its introduction to and use of Python are completely self-contained. Even if you do have some programming experience, the nature of Python and the book's presentation of technical matter won't necessarily relate directly to anything you've learned before: you too might find much to explore here.

The book also assumes no particular knowledge of or experience in bioinformatics or any of the scientific fields to which it relates. It uses real examples from real biological data, and while nearly all of the topics should be familiar to anyone working in the field, there's nothing conceptually daunting about them. Fundamentally, the goal here is to teach you how to write programs that manipulate data.

This book was written with several audiences in mind:

- · Life scientists
- Life sciences students, both undergraduate and graduate
- Technical staff supporting life science research
- Software developers interested in the use of Python in the life sciences

To each of these groups, I offer an introductory message:

Scientists

Presumably you are reading this book because you've found yourself doing, or wanting to do, some programming to support your work, but you lack the computer science or software engineering background to do it as well as you'd like. The book's introduction to Python programming is straightforward, and its

examples are drawn from bioinformatics. You should find the book readable even if you are just curious about programming and don't plan to do any yourself.

Students

This book could serve as a textbook for a one-semester course in bioinformatics programming or an equivalent independent study effort. If you are majoring in a life science, the technical competence you can gain from this book will enable you to make significant contributions to the projects in which you participate. If you are majoring in computer science or software engineering but are intrigued by bioinformatics, this book will give you an opportunity to apply your technical education in that field. In any case, nothing in the book should be intimidating to any student with a basic background either in one of the life sciences or in computing.

Technical staff

You're probably already doing some work managing and manipulating data in support of life science research and development, and you may be accustomed to writing small scripts and performing system maintenance tasks. Perhaps you're frustrated by the limits of your knowledge of computing techniques. Regardless, you have developed an interest in the science and technology of bioinformatics. You want to learn more about those fields and develop your skills in working with biological data. Whatever your training and responsibilities, you should find this book both approachable and helpful.

Programmers

Bioinformatics software differs from most other software in important, though hard to pin down, ways. Python also differs from other programming languages in ways that you will probably find intriguing. This book moves quickly into significant technical material—it does not follow the pattern of a traditional kind of "Programming in..." or "Learning..." or "Introduction to..." book. Though it makes no attempt to provide a bioinformatics primer, the book includes sufficient examples and explanations to intrigue programmers curious about the field and its unusual software needs.



I would like to point out to computer scientists and experienced software developers who may read this book that some very particular choices were made for the purposes of presentation to its intended audience. At the risk of sounding arrogant, I assure you that these are backed by deep theoretical knowledge, extensive experience, and a full awareness of alternatives. These choices were made with the intention of simplifying technical vocabulary and presenting as clear and uniform a view of Python programming as possible. They also were based on the assumption that most people making use of what they learn in this book will not move on to more advanced programming or large-scale software development.

Some things that will appear strange to anyone with significant programming experience are in reality true to a pure "Pythonic" approach. It is delightful to have the opportunity to write in this vocabulary without the need to accommodate more traditional terminology.

The most significant example of this is that the word "variable" is never used in the context of assignment statements or function calls. Python does not assign values to variables in the way that traditional "values in a box" languages do. Instead, like some of the languages that influenced its design, what Python does is assign names to values. The assignment statement should be read from left to right as assigning a name to an existing value. This is a very real distinction that goes beyond the ways languages such as Java and C++ refer to objects through pointer-valued variables.

Another aspect of the book's heavily Pythonic approach is its routine use of comprehensions. Approached by someone familiar with other languages, these can appear quite mysterious. For someone learning Python as a first language, though, they can be more natural and easier to use than the corresponding combinations of assignments, tests, and loops or iterations.

Python

This section introduces the Python language and gives instructions for installing and running Python on your machine.

Some Context

There are many kinds of programming languages, with different purposes, styles, intended uses, etc. Professional programmers often spend large portions of their careers working with a single language, or perhaps a few similar ones. As a result, they are often unaware of the many ways and levels at which programming languages can differ. For educational and professional development purposes, it can be extremely valuable for programmers to encounter languages that are fundamentally different from the ones with which they are familiar.

The effects of such an encounter are similar to learning a foreign human language from a different culture or language family. Learning Portuguese when you know Spanish is not much of a mental stretch. Learning Russian when you are a native English speaker is. Similarly, learning Java is quite easy for experienced C++ programmers, but learning Lisp, Smalltalk, ML, or Perl would be a completely different experience.

Broadly speaking, programming languages embody combinations of four paradigms. Some were designed with the intention of staying within the bounds of just one, or perhaps two. Others mix multiple paradigms, although in these cases one is usually dominant. The paradigms are:

Procedural

This is the traditional kind of programming language in which computation is described as a series of steps to be executed by the computer, along with a few mechanisms for branching, repetition, and subroutine calling. It dates back to the earliest days of computing and is still a core aspect of most modern languages, including those designed for other paradigms.

Declarative

Declarative programming is based on statements of facts and logical deduction systems that derive further facts from those. The primary embodiment of the logic programming paradigm is Prolog, a language used fairly widely in Artificial Intelligence (AI) research and applications starting in the 1980s. As a purely logic-based language, Prolog expresses computation as a series of predicate calculus assertions, in effect creating a puzzle for the system to solve.

Functional

In a purely functional language, all computation is expressed as function calls. In a truly pure language there aren't even any variable assignments, just function parameters. Lisp was the earliest functional programming language, dating back to 1958. Its name is an acronym for "LISt Processing language," a reference to the kind of data structure on which it is based.

Lisp became the dominant language of AI in the 1960s and still plays a major role in AI research and applications. The language has evolved substantially from its early beginnings and spawned many implementations and dialects, although most of these disappeared as hardware platforms and operating systems became more standardized in the 1980s.

A huge standardization effort combining ideas from several major dialects and a great many extensions, including a complete object-oriented (see below) component, was undertaken in the late 1980s. This effort resulted in the now-dominant CommonLisp.* Two important dialects with long histories and extensive current use are Scheme and Emacs Lisp, the scripting language for the Emacs editor. Other functional programming languages in current use are ML and Haskell.

Object-oriented

Object-oriented programming was invented in the late 1960s, developed in the research community in the 1970s, and incorporated into languages that spread widely into both academic and commercial environments in the 1980s (primarily Smalltalk, Objective-C, and C++). In the 1990s this paradigm became a key part of modern software development approaches. Smalltalk and Lisp continued to be used, C++ became dominant, and Java was introduced. Mac OS X, though built on a Unix-like kernel, uses Objective-C for upper layers of the system, especially the user interface, as do applications built for Mac OS X. JavaScript, used primarily to program web browser actions, is another object-oriented language. Once a

^{*} See http://www.lispworks.com/documentation/HyperSpec/Body/01_ab.htm.

radical innovation, object-oriented programming is today very much a mainstream paradigm.

Another dimension that distinguishes programming languages is their primary intended use. There have been languages focused on string matching, languages designed for embedded devices, languages meant to be easy to learn, languages built for efficient execution, languages designed for portability, languages that could be used interactively, languages based largely on list data structures, and many other kinds.

Language designers, whether consciously or not, make choices in these and other dimensions. Subsequent evolutions of their languages are subject to market forces, intellectual trends, hardware developments, and so on. These influences may help a language mature and reach a wider audience. They may also steer the language in directions somewhat different from those originally intended.

The Python Language

Simply put, Python is a beautiful language. It is effective for everything from teaching new programmers to advanced computer science study, from simple scripts to sophisticated advanced applications. It has always had some purchase in bioinformatics, and in recent years its popularity has been increasing rapidly. One goal of this book is to help significantly expand Python's use for bioinformatics programming.

Python features a syntax in which the ends of statements are marked only by the end of a line, and statements that form part of a compound statement are indented relative to the lines of code that introduce them. The semicolons or keywords that end statements and the braces that group statements in other languages are entirely absent.

Programmers familiar with "standard syntax" languages often find Python's uncluttered syntax deeply disconcerting. New programmers have no such problem, and for them, this simple and readable syntax is far easier to deal with than the visually arcane constructions using punctuation (with the attendant compilation errors that must be confronted). Traditional programmers should reconsider Python's syntax after performing this experiment:

- 1. Open a file containing some well-formatted code.
- 2. Delete all semicolons, braces, and terminal keywords such as end, endif, etc.
- 3. Look at the result.

To the human eye, the simplified code is easier to read—and it looks an awful lot like Python. It turns out that the semicolons, terminal keywords, and braces are primarily for the benefit of the compiler. They are not really necessary for human writers and readers of program code. Python frees the programmer from the drudgery of serving as a compiler assistant.

Python is an interesting and powerful language with respect to computing paradigms. Its skeleton is procedural, and it has been significantly influenced by functional

programming, but it has evolved into a fundamentally object-oriented language. (There is no declarative programming component—of the four paradigms, declarative programming is the one least amenable to fitting together with another.) Few, if any, other languages provide a blend like this as seamlessly and elegantly as does Python.

Installing Python

This book uses Python 3, the language's first non-backward-compatible release. With a few minor changes, noted where applicable, Python 2.x will work for most of the book's examples. There are a few notes about Python 2 in Chapters 1, 3, and 5; they are there not just to help you if you find yourself using Python 2 for some work, but also for when you read Python 2 code. The major exception is that print was a statement in Python 2 but is now a function, allowing for more flexibility. Also, Python 3 reorganized and renamed some of its library modules and their contents, so using Python 2.x with examples that demonstrate the use of certain modules would involve more than a few minor changes.

Determing Which Version of Python Is Installed

Some version of Python 2 is probably installed on your computer, unless you are using Windows. Typing the following into a command-line window (using % as an example of a command-line prompt) will tell you which version of Python is installed as the program called python:

% python -V

The name of the executable for Python 3 may be python3 instead of just python. You can type this:

% python3 -V

to see if that is the case.

If you are running Python in an integrated development environment—in particular IDLE, which is part of the Python installation—type the following at the prompt (>>>) of its interactive shell window to get information about its version:

```
>>> from sys import version
>>> version
```

If this shows a version earlier than 3, look for another version of the IDE on your computer, or install one that uses Python 3. (The Python installation process installs the GUI-based IDLE for whatever version of Python is being installed.)

The current release of Python can be downloaded from http://python.org/download/. Installers are available for OS X and Windows. With most distributions of Linux, you should be able to install Python through the usual package mechanisms. (Get help from someone who knows how to do that if you don't.) You can also download the source,

unpack the archive, and, following the steps in the "Build Instructions" section of the *README* file it contains, configure, "make," and install the software.



If you are installing Python from its source code, you may need to download, configure, make, and install several libraries that Python uses if available. At the end of the "make" process, a list of missing optional libraries is printed. It is not necessary to obtain all the libraries. The ones you'll want to have are:

- curses
- gdbm
- sqlite3[†]
- Tcl/Tk[‡]
- readline

All of these should be available through standard package installers.

Running Python

You can start Python in one of two ways:

- 1. Type python3 on the command line.
- 2. Run an IDE. Python comes with one called IDLE, which is sufficient for the work you'll do in this book and is a good place to start even if you eventually decide to move on to a more sophisticated IDE.

The term *Unix* in this book refers to all flavors thereof, including Linux and Mac OS X. The term *command line* refers to where you type commands to a "shell"—in particular, a Unix shell such as tcsh or bash or a Windows command window—as opposed to typing to the Python interpreter. The term *interpreter* may refer to either the interpreter running in a shell, the "Python Shell" window in IDLE, or the corresponding window in whatever other development environment you might be using.

- † You can find precompiled binaries for most platforms at http://sqlite.org/download.html.
- ‡ See http://www.activestate.com/activetcl.
- § On OS X, a command-line shell is obtained by running the Terminal application, found in the *Utilities* folder in the *Applications* folder. On most versions of Windows, a "Command Prompt" window can be opened either by selecting Run from the Start menu and typing cmd or by selecting Accessories from the Start menu, then the Command Prompt entry of that menu. You may also find an Open Command Line Here entry when you right-click on a folder in a Windows Explorer window; this is perhaps the best way to start a command-line Python interpreter in Windows because it starts Python with the selected folder as the current directory. You may have to change your path settings to include the directory that contains the Python executable file. On a Unix-based system, you do that in the "rc" file of the shell you are using (e.g., ~/.bashrc). On Windows, you need to set the path as an environment variable, a rather arcane procedure that differs among different versions of Windows. You can also type the full path to the Python executable; on Windows, for example, that would probably be C:\python3.1\python.exe.

When Python starts interactively, it prints some information about its version. Then it repeats a cycle in which it:

- 1. Prints the *prompt* >>> to indicate that it is waiting for you to type something
- 2. Reads what you type
- 3. Interprets its meaning to obtain a value
- 4. Prints that value

Throughout the book, the appearance of the >>> prompt in examples indicates the use of the interpreter. Like nearly all command-line interactive applications, the Python interpreter won't pay any attention to what you've typed until you press the Return (Enter) key. Pressing the Return key does not always provide complete input for Python to process, though; as you'll see, it is not unusual to enter multiline inputs. In the command-line interpreter, Python will indicate that it is still waiting for you to complete your input by starting lines following the >>> prompt with IDLE, unfortunately, gives no such indication.

Both IDLE and the command-line interpreter provide simple keyboard shortcuts for editing the current line before pressing Return. There are also keys to recall previous inputs. IDLE provides quite a few additional keyboard shortcuts that are worth learning early on. In addition, if you are using an IDE—IDLE, in particular—you'll be able to use the mouse to click to move the cursor on the input line.

To get more information about using IDLE, select "IDLE Help" from its Help menu. That won't show you the keyboard shortcuts, though; they are listed in the "Keys" tab of IDLE's preferences dialog. Note that you can use that dialog to change the keystroke assignments, as well as the fonts, colors, window size, and so on.

When you want to quit a command-line Python interpreter, simply type Ctrl-D in Unix (including Linux and the Mac OS X Terminal application). In Windows, type Ctrl-Z. You exit an IDE with the usual Quit menu command.

Notes

I end this preface with some notes about things I think will help you make the most of your experience with this book.

Reading and Reference Recommendations

The documentation that comes with the Python installation is excellent, extensive, and well organized, but it can be overwhelming to newcomers. Moreover, the topics this book presents and the way it presents them are designed specifically with bioinformatics students and professionals in mind (though of course it's hoped that it will be valuable to a much wider audience than that). Unless you find yourself needing more information about the Python language or library than is provided in this book while you're reading it, it's probably best to wait until you finish it before spending much time with the documentation. The documentation is aimed primarily at Python programmers. You'll be one when you finish this book, at which point you'll use the documentation all the time.

With respect to the bioinformatics side of things, I trust you won't encounter anything unfamiliar here. But if you do, or you want to delve deeper, Wikipedia is a remarkably deep resource for bioinformatics—for programming and computer science too, for that matter. There are two astoundingly extensive bioinformatics references you should at least have access to, if not actually own:

- Bioinformatics and Functional Genomics, Second Edition, by Jonathan Pevsner (Wiley-Blackwell)
- Bioinformatics: Sequence and Genome Analysis, Second Edition, by David W. Mount (Cold Spring Harbor Laboratory Press)

An unusual collection of essays containing detailed information about approaches to analyzing bioinformatics data and the use of a vast array of online resources and tools is:

• Bioinformatics for Geneticists: A Bioinformatics Primer for the Analysis of Genetic Data, Second Edition, by Michael R. Barnes (Ed.) (Wiley)

Example Code

All the code for this book's examples, additional code, some lists of URLs, data for the examples, and so forth are found on the book's website. In many cases, there is a sequence of files for the same example or set of examples that shows the evolution of the example from its simplest start to where it ends up. In a few cases, there are versions of the code that go beyond what is shown in the book. There are also some examples that are not in the book at all, as well as exercises for each chapter.

Within the book's code examples, statement keywords are in boldface. Comments and documentation are in serif typeface. Some examples use oblique monospace to indicate descriptive "pseudocode" meant to be replaced with actual Python code. A shaded background indicates either code that has changed from the previous version of an example or code that implements a point made in the preceding paragraph(s).

Unfortunate and Unavoidable Vocabulary Overlap

This book's vocabulary is drawn from three domains that, for the most part, are independent of each other: computer science (including data structures and programming language concepts), Python (which has its own names for the program and data structures it offers), and biology. Certain words appear in two or even all three of these domains, often playing a major role and having a different meaning in each. The result is unfortunate and unavoidable collisions of vocabulary. Care was taken to establish

sufficient context for the use of these words to make their meanings clear, but the reader should be prepared for the occasional mental backtrack to try another meaning for a term while attempting to make sense of what is being said.

Even within a single domain, a term's importance does not necessarily rescue it from ambiguity. Consider the almost unrelated meanings of the term "frame" as the offset from the start of a DNA sequence and in the phrase "open reading frame." There can be many open reading frames in a frame and many frames with open reading frames. And sometimes there are three frames to consider, and sometimes also the reverse complement frames, making six. Open reading frames can appear in any of the six reading frames.

The vocabulary overlap is so omnipresent that it is almost humorous. Then again, the words involved are fine words that have meanings in a great many other domains too, so we should not be surprised to encounter them in our three. Even though you have not yet been properly introduced to them, Table P-1 lists some of the most vexing examples. Stay particularly alert when you encounter these, especially when you see the words in code examples.

Table P-1. Domain-ambiguous terms

Term	Biology	Programming	Python
Sequence	Part of a DNA or RNA mole- cule; more often refers to the abstraction thereof, as rep- resented with letters	(Usually) one of a number of data structures that arrange their elements linearly	A linear, and therefore numerically indexable, collection of values
Base	A single nucleotide in a DNA or RNA molecule	Base 10, 16, 2, etc.	Base 10, 16, 2, etc., as used in input and output operations
String	A series of letters represent- ing a DNA, RNA, or amino acid sequence	A sequence of characters, often a "primitive type" of a language	An immutable sequence type named str
Expression	The production of proteins under the control of cellular machinery influenced by life stage, the organ containing the cell, internal states (disease, hunger), and external conditions (dryness, heat)	(1) (Generally) a combination of primitive values, operators, and function calls, with specifics differing significantly among languages	(1) A combination of primitive values, operators, and function calls
		(2) Regular expression: a pattern describ-	(2a) A regular expression string
		ing a set of strings with notations for types of characters, grouping, repetition, and so on, the details of which differ among lan- guages and editors	(2b) A regular expression string compiled into a regular expression object
Туре	The specimen of an organ- ismfirst used to describe and name it	A theoretical construct defined differently in different contexts and implemented differently by different programming lan-	Synonymous with "class," but often used in the context of Python's built-in types, as op-

Term	Biology	Programming	Python
		guages; corresponds roughly to "the kind of thing" something is and "the kind of operations" it supports	posed to classes defined in a Python or externally obtained library or in user code
Translate, translate	Convert DNA codons (base triples) to amino acids according to the genetic code of the organism	Convert computer code in one language into computer code in another, typically lower-level, language	A method of str that uses a table to produce a new str with all the characters of the original replaced by the corresponding entries in the table
Class	One of the levels in the standard taxonomic classification of organisms	In languages that support object- oriented programming, the encapsulated definition of data and related code	As in programming; more spe- cifically, the type of an object, which itself is an object that defines the methods for its instances
Loop	A property of RNA secondary structures (among other meanings)	An action performed repeatedly until some condition is no longer true	An action performed repeatedly until some condition is no longer true
Library	A collection of related sequences, most commonly used in the context of a library of expressed RNA in cDNA form	Like a program, but meant to be used by other programs rather than as a free- standing application; most languages use a core set of libraries and provide a large selection of optional ones	A collection of modules, each containing a collection of related definitions, as in "Python comes with an extensive library of optional tools and facilities"
Complement	The nucleotide with which another always pairs	"Two's complement" is the standard representation of negative integers	
R.E.	Restriction enzyme		Regular expression

Fortunately, while the term "sequence" has a conceptual meaning in Python, there is nothing defined in the language by that name, so we can use it in our descriptions and code examples. Likewise, the name of the string type is str, so we can use the term "string" in descriptions and examples. The lack of overlap in these instances saves a fair amount of awkward clarification that would otherwise be required.

Comments

Write code as you read: it will really help you understand and benefit from what you are reading. Read through the code examples. Look for more detailed code, additional examples, and exercises on the book's website.

Bioinformatics is a fascinating field. Python is a wonderful language. Programming is an exciting challenge. Technologies covered here are important. This book is an invitation to investigate, experience, and learn (more) about all of these topics. Enjoy!

Conventions Used in This Book

The following typographical conventions are used in this book:

Italic

Indicates new terms, URLs, filenames, and file extensions, and is used for documentation and comments in code examples

Constant width

Used for names of values, functions, methods, classes, and other code elements, as well as the code in examples, the contents of files, and printed output

Constant width bold

Indicates user input to be typed at an interactive prompt, and highlights Python statement keywords in examples

Constant width italic

Shows text that should be replaced with user-supplied values; also used in examples for "pseudocode" meant to be replaced with actual Python code



This icon signifies a tip, suggestion, or general note.



This icon indicates a warning or caution.

Some chapters include text and code in specially labeled sidebars. These capture explanations and examples in an abstract form. They are meant to serve as aids to learning the first time you go through the material, and useful references later. There are three kinds of these special sidebars:

STATEMENT

Every kind of Python statement is explained in text and examples and summarized in a statement box.

SQL

SQL boxes are used in Chapter 10 to summarize statements from the database language SQL.

TEMPLATE

In place of theoretical discussions of the many ways a certain kind of code can be written, this book presents generalized abstract patterns that show how such code would typically appear in bioinformatics programs. The idea is that, at least in the early stages of using what you learn in this book, you can use the templates in your programs, replacing the abstract parts with the concrete details of your code.

We'd Like to Hear from You

Every example in this book has been tested, but occasionally you may encounter problems. Mistakes and oversights can occur, and we will gratefully receive details of any that you find, as well as any suggestions you would like to make for future editions. You can contact the author and editor at:

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I thank my students at Northeastern University's Professional Masters in Bioinformatics program for their patience, suggestions, and error detection while using earlier versions of the book's text and code. Special thanks go to Jyotsna Guleria, a graduate of that program, who wrote test programs for the example code that uncovered significant latent errors. (Extended versions of the test programs can be found on the book's website.) Finally, I hope what I have produced justifies what my friends, family, and colleagues endured during its creation—especially Janet, my wife, whose unwavering support during the book's writing made the project possible.



Primitives

Computer programs manipulate data. This chapter describes the simplest kinds of Python data and the simplest ways of manipulating them. An individual item of data is called a *value*. Every value in Python has a *type* that identifies the *kind* of value it is. For example, the type of 2 is int. You'll get more comfortable with the concepts of types and values as you see more examples.

The Preface pointed out that Python is a multiparadigm programming language. The terms "type" and "value" come from traditional procedural programming. The equivalent object-oriented terms are *class* and *object*. We'll mostly use the terms "type" and "value" early on, then gradually shift to using "class" and "object" more frequently. Although Python's history is tied more to object-oriented programming than to traditional programming, we'll use the term *instance* with both terminologies: each value is an instance of a particular type, and each object is an instance of a particular class.

Simple Values

Types for some simple kinds of values are an integral part of Python's implementation. Four of these are used far more frequently than others: *logical* (Boolean), *integer*, *float*, and *string*. There is also a special no-value value called None.

When you enter a value in the Python interpreter, it prints it on the following line:

```
>>> 90
90
```

When the value is None, nothing is printed, since None means "nothing":

```
>>> None >>>
```

If you type something Python finds unacceptable in some way, you will see a multiline message describing the problem. Most of what this message says won't make sense

until we've covered some other topics, but the last line should be easy to understand and you should learn to pay attention to it. For example:

```
Traceback (most recent call last):
 File "<pyshell#7>", line 1, in <module>
NameError: name 'Non' is not defined
```

When a # symbol appears on a line of code, Python ignores it and the rest of the line. Text following the # is called a *comment*. Typically comments offer information about the code to aid the reader, but they can include many other kinds of text: a programmer's notes to fix or investigate something, a reference (documentation entry, book title, URL, etc.), and so on. They can even be used to "comment out" code lines that are not working or are obsolete but still of interest. The code examples that follow include occasional comments that point out important details.

Booleans

There are only two Boolean values: True and False. Their type is bool. Python names are "case-sensitive," so true is not the same as True:

```
>>> True
True
>>> False
False
```

Integers

There's not much to say about Python integers. Their type is int, and they can have as many digits as you want. They may be preceded by a plus or minus sign. Separators such as commas or periods are not used:

```
>>> 14
14
>>> -1
-1
>>> 1112223334445556667778889990000000000000
                                                       # a very large integer!
1112223334445556667778889990000000000000
```



Python 2: A distinction is made between integers that fit within a certain (large) range and those that are larger; the latter are a separate type called long.

Integers can also be entered in hexadecimal notation, which uses base 16 instead of base 10. The letters A through F represent the hexadecimal digits 10 through 15. Hexadecimal notation begins with **ox**. For example:

```
\# (1 \times 16) + 2
>>> 0x12
18
>>> 0xA40
                                            \# (10 \times 16 \times 16) + (4 \times 16) + 0
2624
>>> 0xFF
                                            #(15 \times 16) + 15
255
```

The result of entering a hexadecimal number is still an integer—the only difference is in how you write it. Hexadecimal notation is used in a lot of computer-related contexts because each hexadecimal digit occupies one half-byte. For instance, colors on a web page can be specified as a set of three one-byte values indicating the red, green, and blue levels, such as FFA040.

Floats

"Float" is an abbreviated version of the term "floating point," which refers to a number that is represented in computer hardware in the equivalent of scientific notation. Such numbers consist of two parts: digits and an exponent. The exponent is adjusted so the decimal point "floats" to just after the first digit (or just before, depending on the implementation), as in scientific notation.

The written form of a float always contains a decimal point and at least one digit after it:

```
>>> 2.5
2.5
```

You might occasionally see floats represented in a form of scientific notation, with the letter "e" separating the base from the exponent. When Python prints a number in scientific notation it will always have a single digit before the decimal point, some number of digits following the decimal point, a + or - following the e, and finally an integer. On input, there can be more than one digit before the decimal point. Regardless of the form used when entering a float, Python will output very small and very large numbers using scientific notation. (The exact cutoffs are dependent on the Python implementation.) Here are some examples:

```
>>> 2e4
                                     # Scientific notation, but...
20000.0
                                     # within the range of ordinary floats.
>>> 2e-2
0.02
>>>.0001
                                     # Within the range of ordinary floats
0.0001
                                     # so printed as an ordinary float.
>>>.00001
                                     # An innocent-looking float that is
                                     # smaller than the lower limit, so e.
1e-05
                                     # A float with many digits that is
>>> 1002003004005000.
1002003004005000.0
                                     # smaller than the upper limit, so no e.
>>> 100200300400500060.
                                     # Finally, a float that is larger than the
1.0020030040050006e+17
                                     # upper limit, so printed with an e.
```

Strings

Strings are series of Unicode* characters. Their type is str. Many languages have a separate "character" type, but Python does not: a lone character is simply a string of length one. A string is enclosed in a pair of single or double quotes. Other than style preference, the main reason to choose one or the other kind of quote is to make it convenient to include the other kind inside a string.

If you want a string to span multiple lines, you must enclose it in a matched pair of three single or double quotes. Adding a backslash in front of certain characters causes those characters to be treated specially; in particular, '\n' represents a line break and '\t' represents a tab.



Python 2: Strings are composed of one-byte characters, not Unicode characters; there is a separate string type for Unicode, designated by preceding the string's opening quote with the character u.

We will be working with strings a lot throughout this book, especially in representing DNA/RNA base and amino acid sequences. Here are the amino acid sequences for some unusually small bacterial restriction enzymes:[†]

>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'

MNKMDLVADVAEKTDLSKAKATEVIDAVFA

 $\gt\gt\gt\gt$ "AARHQGRGAPCGESFWHWALGADGGHGHAQPPFRSSRLIGAERQPTSDCRQSLQQSPPC"

AARHQGRGAPCGESFWHWALGADGGHGHAQPPFRSSRLIGAERQPTSDCRQSLQQSPPC

>>> """MKQLNFYKKN SLNNVQEVFS YFMETMISTN RTWEYFINWD KVFNGADKYR NELMKLNSLC GS LFPGEELK SLLKKTPDVV KAFPLLLAVR DESISLLD"""

'MKQLNFYKKN SLNNVQEVFS YFMETMISTN RTWEYFINWD KVFNGADKYR NELMKLNSLC GS LFPGEELK\nSLLKKT PDVV KAFPLLLAVR DESISLLD'

>>> '''MWNSNLPKPN AIYVYGVANA NITFFKGSDI LSYETREVLL KYFDILDKDE RSLKNALKD LEN PFGFAPYI RKAYEHKRNF LTTTRLKASF RPTTF'''

'MWNSNLPKPN AIYVYGVANA NITFFKGSDI LSYETREVLL KYFDILDKDE RSLKNALKDL EN\nPFGF APYI RKAYEHKRNF LTTTRLKASF RPTTF'

There are three situations that cause input or output to begin on a new line:

- You hit Return as you are typing inside a triple-quoted string.
- You keep typing characters until they "wrap around" to the next line before you press Return.
- The interpreter responds with a string that is too long to fit on one line.

^{*} Unicode characters occupy between one and four bytes each in memory, depending on several factors. See http://docs.python.org/3.1/howto/unicode.html for details (in particular, http://docs.python.org/3.1/howto/unicode.html for general information about Unicode outside of Python, consult http://www.unicode.org/standard/WhatIsUnicode.html, http://www.unicode.org/standard/principles.html, and http://www.unicode.org/resources.

[†] Data for these examples was obtained from the "Official REBASE Homepage" site. Files in formats used by various applications can be downloaded from http://rebase.neb.com/rebase/rebase.files.html.

Only the first one is "real." The other two are simply the effect of output "line wrapping" like what you would see in text editors or email programs. In the second and third situations, if you change the width of the window the input and output strings will be "rewrapped" to fit the new width. The first case does *not* cause a corresponding line break when the interpreter prints the string—the Return you typed becomes a '\n' in the string.

Normally, Python uses a pair of single quotes to enclose strings it prints. However, if the string contains single quotes (and no double quotes), it will use double quotes. It never prints strings using triple quotes; instead, the line breaks typed inside the string become '\n's.

Expressions

An *operator* is a symbol that indicates a calculation using one or more *operands*. The combination of the operator and its operand(s) is an *expression*.

Numeric Operators

A unary operator is one that is followed by a single operand. A binary operator is one that appears between two operands. It isn't necessary to surround operators with spaces, but it is good style to do so. Incidentally, when used in a numeric expression, False is treated as 0 and True as 1.

Plus and minus can be used as either unary or binary operators:

```
>>> -1
                      # unary minus
-1
>>> 4 + 2
>>> 4 - 1
>>> 4 * 3
12
```

The power operator is ** (i.e., n^k is written n ** k):

```
>>> 2 ** 10
1024
```

There are three operators for the division of one integer by another: / produces a float, // (floor division) an integer with the remainder ignored, and % (modulo) the remainder of the floor division. The formal definition of floor division is "the largest integer not greater than the result of the division":

```
>>> 11 / 4
2.75
>>> 11 // 4
                        # "floor" division
```

```
>>> 11 % 4
                         # remainder of 11 // 3
```



Python 2: The / operator performs floor division when both operands are ints, but ordinary division if one or both operands are floats.

Whenever one or both of the operators in an arithmetic expression is a float, the result will be a float:

```
>>> 2.0 + 1
3.0
>>> 12 * 2.5
30.0
>>> 7.5 // 2
3.0
```



While the value of floor division is *equal* to an integer value, its *type* may not be integer! If both operands are ints, the result will be an int, but if either or both are floats, the result will be a float that represents an integer.

The result of an operation does not always print the way you might expect. Consider the following numbers:

```
>>> .009
.009
>>> .01
.01
>>> .029
.029
>>> .03
.03
>>> .001
.001
```

So far, everything is as expected. If we subtract the first from the second and the third from the fourth, we should in both cases get the result .001. Typing in .001 also gives the expected result. However, typing in the subtraction operations does not:

```
>>> .03 - .029
0.00099999999999974
>>> .01 - .009
0.0010000000000000009
```

Strange results like this arise from two sources:

 For a given base, only some rational numbers have "exact" representations—that is, their decimal-point representations terminate after a finite number of digits. The rest end in an infinitely repeating sequence of digits (e.g., 1/3 = 0.3333333...).

 A computer stores rational numbers in a finite number of binary digits; the binary representation of a rational number may in fact have an exact binary representation, but one that would require more digits than are used.



A rational number is one that can be expressed as a/b, where b is not zero; the decimal-point expression of a rational number in a given number system either has a finite number of digits or ends with an infinitely repeating sequence of digits. There's nothing wrong with the binary system: whatever base is used, some real numbers have exact representations and others don't. Just as only some rational numbers have exact decimal representations, only some rational numbers have exact binary representations.

As you can see from the results of the two division operations, the difference between the ideal rational number and its actual representation is quite small, but in certain kinds of computations the differences do accumulate.[‡]

Here's an early lesson in an extremely important programming principle: don't trust what you see! Everything printed in a computing environment or by a programming language is an interpretation of an internal representation. That internal representation may be manipulated in ways that are intended to be helpful but can be misleading. In the preceding example, 0.009 in fact does not have an exact binary representation. In Python 2, it would have printed as 0.008999999999993, and 0.003 would have printed as 0.008999999999999993. The difference is that Python 3 implements a more sophisticated printing mechanism for rational numbers that makes some of them look as they would have had you typed them.

Logical Operations

Python, like other programming languages, provides operations on "truth values." These follow the mathematical laws of Boolean logic. The classic Boolean operators are not, and, and or. In Python, those are written just that way rather than using special symbols:

```
>>> not True
False
>>> not False
True
>>> True and True
True
>>> True and False
False
>>> True or True
True
```

[‡] A computer science field called "numerical analysis" provides techniques for managing the accumulation of such errors in complex or repetitive computations.

```
>>> True or False
>>> False and False
False
>>> False or True
```



The results of and and or operations are not converted to Booleans. For and expressions, the first operand is returned if it is false; otherwise, the second operand is returned. For or expressions, the first operand is returned if it is true; otherwise, the second operand is returned. For example:

```
>>> '' and 'A'
                             # Not False: " is a false value
                             # Read as (0 and 1) or 2
>>> 0 and 1 or 2
                             # Not True: 2 is a false value
```

While confusing, this can be useful; we'll see some examples later.

The operands of and and or can actually be anything. None, 0, 0.0, and the empty string, as well as the other kinds of "empty" values explained in Chapter 3, are considered False. Everything else is treated as True.



To avoid repetition and awkward phrases, this book will use "true" and "false" in regular typeface to indicate values considered to be True and False, respectively. It will only use the capitalized words True and False in the code typeface when referring to those specific Boolean values.

There is one more logical operation in Python that forms a conditional expression. Written using the keywords if and else, it returns the value following the if when the condition is true and the value following the else when it is false. We'll look at some more meaningful examples a bit later, but here are a few trivial examples that show what conditional expressions look like:

```
>>> 'yes' if 2 - 1 else 'no'
'yes'
>>> 'no' if 1 % 2 else 'no'
```

In addition to the Boolean operators, there are six comparison operators that return Boolean values: ==, !=, <, <=, >, and >=. These work with many different kinds of operands:

```
>>> 2 == 5 // 2
True
>>> 3 > 13 % 5
False
>>> 'one' < 'two'
True
```

```
>>> 'one' != 'one'
```

You may already be familiar with logical and comparison operations from other computer work you've done, if only entering spreadsheet formulas. If these are new to you, spend some time experimenting with them in the Python interpreter until you become comfortable with them. You will use them frequently in code you write.

String Operations

There are four binary operators that act on strings: in, not in, +, and *. The first three expect both operands to be strings. The last requires the other operator to be an integer. A one-character substring can be extracted with *subscription* and a longer substring by *slicing*. Both use square brackets, as we'll see shortly.

String operators

The in and not in operators test whether the first string is a substring of the second one (starting at any position). The result is True or False:

```
>>> 'TATA' in 'TATATATATATATATATATATATA'
True
>>> 'AA' in 'TATATATATATATATATATATATA'
>>> 'AA' not in 'TATATATATATATATATATATATA'
True
```

A new string can be produced by *concatenating* two existing strings. The result is a string consisting of all the characters of the first operand followed by all the characters of the second. Concatenation is expressed with the plus operator:

```
>>> 'AC' + 'TG'
'ACTG'
>>> 'aaa' + 'ccc' + 'ttt' + 'ggg'
'aaaccctttggg'
```

A string can be repeated a certain number of times by multiplying it by an integer:

```
>>> 'TA' * 12
'TATATATATATATATATATATA'
>>> 6 * 'TA'
'ΤΑΤΑΤΑΤΑΤΑΤΑ'
```

Subscription

Subscription extracts a one-character substring of a string. Subscription is expressed with a pair of square brackets enclosing an integer-valued expression called an *index*. The first character is at position 0, not 1:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[0]
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[1]
'N'
```

The index can also be negative, in which case the index is counted from the end of the string. The last character is at index -1:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[-1]
'A'
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[-5]
'D'
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[7 // 2]
```

As Figure 1-1 shows, starting at 0 from the beginning or end of a string, an index can be thought of as a label for the character to its right. The end of a string is the position one after the last element. If you are unfamiliar with indexing in programming languages, this is probably an easier way to visualize it than if you picture the indexes as aligned with the characters.

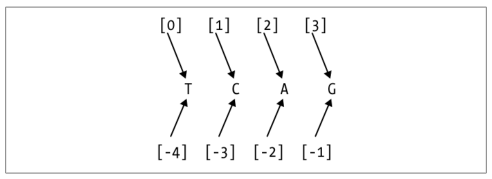


Figure 1-1. Index positions in strings

Attempting to extract a character before the first or after the last causes an error, as shown here:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[50]
Traceback (most recent call last):
   File "<pyshell#14>", line 1, in <module>
     'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[50]
IndexError: string index out of range
```

The last line reports the nature of the error, while the next-to-last line shows the input that caused the error.

Slicing

Slicing extracts a series of characters from a string. You'll use it often to clearly and concisely designate parts of strings. Figure 1-2 illustrates how it works.

The character positions of a slice are specified by two or three integers inside square brackets, separated by colons. The first index indicates the position of the first character to be extracted. The second index indicates where the slice ends. The character at that position is not included in the slice. A slice [m:n] would therefore be read as "from

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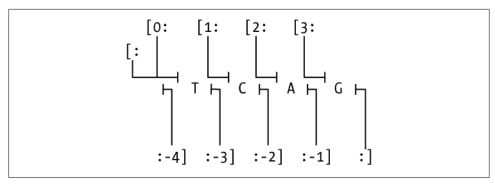


Figure 1-2. String slicing

character m up to but not including character n." (We'll explore the use of the third index momentarily). Here are a few slicing examples:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[1:4]
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[4:-1]
'DLVADVAEKTDLSKAKATEVIDAVF'
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[-5:-4]
```

Either of the indexes can be positive, indicating "from the beginning," or negative, indicating "from the end." If neither of the two numbers is negative, the length of the resulting string is the difference between the second and the first. If either (or both) is negative, just add it to the length of the string to convert it to a nonnegative number.

What if the two numbers are the same? For example:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[5:5]
```

Since this reads as "from character 5 up to but not including character 5," the result is an empty string. Now, what about character positions that are out of order—i.e., where the first character occurs after the second? This results in an empty string too:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[-4:-6]
```

For subscription, the index must designate a character in the string, but the rules for slicing are less constraining.

When the slice includes the beginning or end of the string, that part of the slice notation may be omitted. Note that omitting the second index is not the same as providing -1 as the second index—omitting the second index says to go up to the end of the string, one past the last character, whereas -1 means go up to the penultimate character (i.e., up to but not including the last character):

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[:8]
'MNKMDLVADVAEKTDLSKAKAT'
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[9:]
```

```
'VAEKTDLSKAKATEVIDAVFA'
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[9:-1]
'VAEKTDLSKAKATEVIDAVF'
```

In fact, both indexes can be omitted, in which case the entire string is selected:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[:]
'MNKMDI VADVAFKTDI SKAKATEVTDAVFA'
```

Finally, as mentioned earlier, a slice operation can specify a third number, also following a colon. This indicates a number of characters to skip after each one that is included, known as a step. When the third number is omitted, as it often is, the default is 1, meaning don't skip any. Here's a simple example:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[0:9:3]
'MMV'
```

This example's result was obtained by taking the first, fourth, and seventh characters from the string. The step can be also be a negative integer. When the step is negative, the slice takes characters in reverse order. To get anything other than an empty string when you specify a negative step, the start index must be greater than the stop index:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[16:0:-4]
```

Notice that the first character of the string is not included in this example's results. The character at the stop index is never included. Omitting the second index so that it defaults to the beginning of the string—beginning, not end, because the step is negative—results in a string that does include the first character, assuming the step would select it. Changing the previous example to omit the 0 results in a longer string:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[16::-4]
'SKDDM'
```

Omitting the first index when the step is negative means start from the end of the string:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[:25:-1]
'AFVA'
```

A simple but nonobvious slice expression produces a reversed copy of a string: s[::-1]. This reads as "starting at the end of the string, take every character up to and including the first, in reverse order":

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'[::-1]
'AFVADIVETAKAKSLDTKEAVDAVLDMKNM'
```

Calls

We'll look briefly at calls here, deferring details until later. A call is a kind of expression.

Function calls

The simplest kind of call invokes a function. A call to a function consists of a function name, a pair of parentheses, and zero or more argument expressions separated by commas. The function is *called*, does something, then *returns* a value. Before the function is called the argument expressions are evaluated, and the resulting values are passed to the function to be used as input to the computation it defines. An argument can be any kind of expression whose result has a type acceptable to the function. Those expressions can also include function calls.

Each function specifies the number of arguments it is prepared to receive. Most functions accept a fixed number—possibly zero—of arguments. Some accept a fixed number of required arguments plus some number of optional arguments. We will follow the convention used in the official Python documentation, which encloses optional arguments in square brackets. Some functions can even take an arbitrary number of arguments, which is shown by the use of an ellipsis.

Python has a fairly small number of "built-in" functions. Some of the more frequently used are:

len(ara)

Returns the number of characters in *arg* (although it's actually more general than that, as will be discussed later)

```
print(args...[, sep=seprstr][, end=endstr])
```

Prints the arguments, of which there may be any number, separating each by a seprstr (default ' ') and omitting certain technical details such as the quotes surrounding a string, and ending with an *endstr* (default '\n')



Python 2: print is a statement, not a function. There is no way to specify a separator. The only control over the end is that a final comma suppresses the newline.

input(string)

Prompts the user by printing *string*, reads a line of input typed by the user (which ends when the Return or Enter key is pressed), and returns the line as a string



Python 2: The function's name is raw input.

Here are a few examples:

```
>>> len('TATA')
>>> print('AAT', 'AAC', 'AAG', 'AAA')
AAT AAC AAG AAA
>>> input('Enter a codon: ')
Enter a codon: CGC
'CGC'
>>>
```

Here are some common numeric functions in Python:

```
abs(value)
```

Returns the absolute value of its argument

```
max(args...)
```

Returns the maximum value of its arguments

```
min(args...)
```

Returns the minimum value of its arguments

Types can be called as functions too. They take an argument and return a value of the type called. For example:

```
str(arg)
```

Returns a string representation of its argument

int(ara)

Returns an integer derived from its argument

float(arg)

Returns a float derived from its argument

bool(ara)

Returns False for None, zeros, empty strings, etc., and True otherwise; rarely used, because other types of values are automatically converted to Boolean values wherever Boolean values are expected

Here are some examples of these functions in action:

```
>>> str(len('TATA'))
'4'
>>> int(2.1)
>>> int('44')
>>> bool('')
False
>>> bool(' ')
True
>>> float(3)
3.0
```



Using int is the only way to guarantee that the result of a division is an integer. As noted earlier, // is the floor operator and results in a float if either operand is a float.

There is a built-in help facility for use in the Python interpreter. Until we've explored more of Python, much of what the help functions print will probably appear strange or even unintelligible. Nevertheless, the help facility is a useful tool even at this early stage. You can use either of these commands to access it:

help()

Enters the interactive help facility

help(x)

Prints information about x, which can be anything (a value, a type, a function, etc.); help for a type generally includes a long list of things that are part of the type's implementation but not its general use, indicated by names beginning with underscores

Occasionally your code needs to test whether a value is an instance of a certain type; for example, it may do one thing with strings and another with numbers. You can do this with the following built-in function:

isinstance(x, sometype)

Returns True if x is an instance of the type (class) *sometype*, and False otherwise

Method calls

Many different types of values can be supplied as arguments to Python's built-in functions. Most functions, however, are part of the implementation of a specific type. These are called *methods*. Calling a method is just like calling a function, except that the first argument goes before the function name, followed by a period. For example, the method count returns the number of times its argument appears in the string that precedes it in the call. The following example returns 2 because the string 'DL' appears twice in the longer string:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'.count('DL')
```

Except for having their first argument before the function name, calls to methods have the same features as calls to ordinary functions: optional arguments, indefinite number of arguments, etc. Here are some commonly used methods of the str type:

```
string1.count(string2[, start[, end]])
```

Returns the number of times string2 appears in string1. If start is specified, starts counting at that position in string1; if end is also specified, stops counting before that position in *string1*.

```
string1.find(string2[, start[, end]])
```

Returns the position of the last occurrence of string2 in string1; -1 means string2 was not found in string1. If start is specified, starts searching at that position in string1; if end is also specified, stops searching before that position in string1.

```
string1.startswith(string2[, start[, end]])
```

Returns True or False according to whether string2 starts with string1. If start is specified, uses that as the position at which to start the comparison; if *end* is also specified, stops searching before that position in *string1*.

```
string1.strip([string2])
```

Returns a string with all characters in *string2* removed from its beginning and end; if *string2* is not specified, all whitespace is removed.

```
string1.lstrip([string2])
```

Returns a string with all characters in string2 removed from its beginning; if *string2* is not specified, all whitespace is removed.

```
string1.rstrip([string2])
```

Returns a string with all characters in *string2* removed from its end; if *string2* is not specified, all whitespace is removed.

Here are some examples of method calls in action:

```
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'.find('DL')
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'.find('DL', 5)
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'.find('DL', 5, 12)
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'.startswith('DL')
>>> 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'.startswith('DL', 4)
```

The restriction enzyme with the amino acid sequence in these examples recognizes the site with the base sequence TCCGGA. It's easy enough to find the first location in a DNA base sequence where this occurs:

```
>>> 'AAAAATCCCGAGGCGGCTATATAGGGCTCCGGAGGCGTAATATAAAA'.find('TCCGGA')
27
>>>
```

If the recognition site did not occur in the sequence, find would have returned -1.

Compound Expressions

The examples of operator expressions that we've looked at thus far have had only a single operator. However, just as in traditional algebra, operators can be compounded in a series. For example:

```
>>> 2 * 3 + 4 - 1
```

This is read as "2*3 is 6, 6+4 is 10, and 10-1 is 9." The story isn't quite that simple, though. Consider the following example:

```
>>> 4 + 2 * 3 - 1
9
```

Reading from left to right, we'd have "4+2 is 6, 6*3 is 18, 18-1 is 17," not 9. So why do we get 9 as the result? Programming languages incorporate operator precedence rules that determine the order in which operations in a series should be performed. Like

most programming languages, Python performs multiplications and divisions first and then goes back and performs additions and subtractions.

You can indicate your intended interpretation of a sequence of operations by surrounding parts of an expression with parentheses. Everything inside a pair of parentheses will be evaluated completely before the result is used in another operation. For instance, parentheses could be used as follows to make the result of the preceding example be 17:

```
>>> (4 + 2) * 3 - 1
```

Comparisons can be combined to form "between" expressions:

```
>>> 1 < 4 < 6
True
>>> 2 <= 2 < 5
True
>>> 2 < 2 < 5
False
```

Strings can participate in sequences of operations:

```
>>> 'tc' in ('ttt' + 'ccc' + 'ggg' + 'aaa')
>>> 'tc' in 't' * 3 + 'c' * 3 + 'g' * 3 + 'a' * 3
```

The second variation demonstrates that * has a higher precedence than +, and + has a higher precedence than in. Don't hesitate to use parentheses if you have any doubt about the interpretation of operation series.

Here is a list of the operators mentioned in this chapter, ordered from highest precedence to lowest:

- Calls
- Slicings
- Subscriptions
- Exponentiation (**)
- Unary +, -
- Multiplication, division, and remainder (*, /, //, %)
- Addition and subtraction (+, -)
- Comparisons (==, !=, <, <=, >, >=)
- Membership (in, not in)
- Boolean not (not)
- Boolean and (and)
- Boolean or (or)

Tips, Traps, and Tracebacks

Tips

• Don't trust what you see! Everything printed out in a computing environment or by a programming language is an interpretation of an internal representation. The visible interpretation may not be what you anticipated, even though the internal representation is actually the result you expected.

Statements and expressions

- The results of and and or expressions are not converted to Booleans. For and expressions, the first operand is returned if it is false, and otherwise the second operand is returned. For or expressions, the first operand is returned if it is true, and otherwise the second operand is returned. For example, '' and 'A' evaluates to '', not False, while '' or 'A' evaluates to 'A', not True. We'll see examples later of idioms based on this behavior.
- Function calls are both expressions and statements.
- Experiment with using the sep and end keyword arguments to print. They give you more control over your output. The default is to separate every argument by a space and end with a newline.
- A method call is simply a function call with its first argument moved before the function name, followed by a period.
- If you are ever in doubt about the order in which an expression's operations are performed, use parentheses to indicate the ordering you want. Parentheses can sometimes help make the code more readable. They are never required in operation expressions.

Running Python interactively

- Start the Python interpreter from the command line by typing python at a command prompt. Here are a few points to keep in mind:
 - —If the initial message you see when Python starts indicates that its version number begins with a 2, exit and try typing python3. If that doesn't work, try including a version number (e.g., python3.1 or python3.2).
 - —If that doesn't work, either you don't have Python 3 installed or it's not on the path used in your command-line environment. If you don't know how to add

[§] Command line is a term that refers to an interactive terminal-like window: a Unix shell, OS X Terminal window, or Windows Command window. The command line prompts for input and executes the commands you type.

it, find someone knowledgeable about the command-line environment in your operating system to help you resolve the problem.

- The way to exit Python follows each platform's usual conventions: Ctrl-D on Unixbased systems, Ctrl-Z on Windows variants. You can also type quit().
- In Unix and OS X shells, depending on how Python was installed, you may be able to edit the current line you are typing to Python and navigate back and forth in the history of inputs. After you've typed at least one line to Python, try Ctrl-P or the up arrow. If that changes the input to what you typed previously, the editing capability is functioning. You can use Ctrl-P or the down arrow to move to a later line in the input history. Following are some editing operations that work on the current line:

Ctrl-A

Go to the beginning of the line.

Ctrl-E

Go to the end of the line.

Ctrl-B or left arrow

Move one character to the left.

Ctrl-F or right arrow

Move one character to the right.

Backspace

Delete the preceding character.

Ctrl-D

Delete the next character.

Ctrl-K

Delete the rest of the line after the cursor.

Ctrl-Y

"Yank" the last killed text into the line at the location of the cursor.

Ctrl-_ (underscore)

Undo; can be repeated.

Ctrl-R

Search incrementally for a preceding input line.

Ctrl-S

Search incrementally for a subsequent input line.

If not, the Python you are using was built without the readline system (not Python) library. If you configured, compiled, and installed Python yourself, you probably know how to get the readline library, install it, and repeat the configure-compile-install process. If not, you will have no idea what any of this is about, and there probably isn't anything you can do about it.

Return

Give the current line to the interpreter. Similar functionality may be available when Python is run in a Windows command window.

Traps

- The value of a floor division (//) equals an integer but has the type int only if both operands were ints; otherwise, the value is a float that prints with a 0 after the decimal point.
- The result of an operation with a float operand may produce a result very slightly more or very slightly less than its "true" mathematical equivalent.
- Remember that the first element of a string is at index 0 and the last at -1.
- The index in a string indexing expression must be greater than or equal to 0 and less than the length of the string. (The restriction does not apply to slices.)
- In a function call with more than one argument, every argument except the last must be followed by a comma. Usually omitting a comma will cause syntax errors, but in some situations you will accidentally end up with a syntactically correct expression that is not what you intended.
- Omitting a right parenthesis that closes a function call's argument list results in a syntax error message pointing to the line *after* the one containing the function call.
- Function and method calls with no arguments must still be followed by (an empty) pair of) parentheses. Failing to include them will *not* lead to a syntax error, because the value of the name of the function is the function itself—a legitimate value but it will lead to very unexpected results and, often, runtime errors.

Tracebacks

Representative error messages include:

NameError: 'Non' is not defined

Python doesn't recognize a name (more on this in the next chapter).

IndexError: string index out of range

For a string of length N, an index (i.e., the value between square brackets) must be in the range -N <= index < N-1.

SyntaxError

Python syntax violation.

ZeroDivisionError

/, //, or % with **0** as the second operand.

Names, Functions, and Modules

In this chapter we'll see how to name values, define new functions, and incorporate optional software from the Python library. All of these operations rest on Python mechanisms for naming and interpreting names.

A Python *name* consists of an arbitrary number of letters, underscores, and digits. The only restriction is that the first character must not be a digit (otherwise Python would interpret the digit as the beginning of a number). Names beginning with two underscores are special in one of several ways that will be explained later, so names you choose will almost always begin with a letter.

A name is used to refer to something—a primitive value, a function, or any of a number of other possibilities. *A name is not the same as a string*. A name refers to something, whereas a string has no intrinsic meaning to Python. Unlike strings, names are not enclosed in quotes. Giving a name to a value is called *binding*. One value may have multiple names bound to it, as Figure 2-1 illustrates.

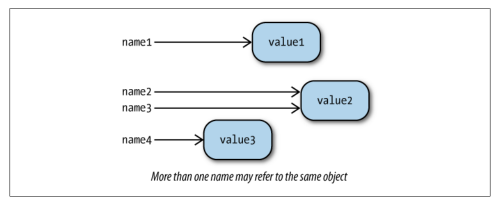


Figure 2-1. Names bound to objects

The same name can mean different things in different contexts. The technical term for such a context is a *namespace*. The types and built-in functions we saw in the previous

chapter are all part of the global namespace. A separate namespace is associated with each type, such as str. This enables each type to have its own version of common methods, such as count and find. Which version gets executed is determined by the type of the value through which the method is called. We'll see more details about all of this later, but Figure 2-2 should give you an idea of how namespaces in Python work.

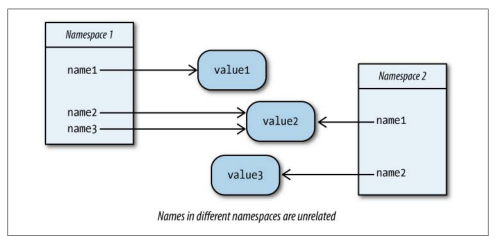


Figure 2-2. Names in different namespaces bound to objects

We've already seen some kinds of things that have names: types such as str, functions such as len, and methods such as str.count. Here's what Python responds with when given these names as input:

```
>>> str
<class 'str'>
>>> len
<built-in function len>
>>> str.count
<method 'count' of 'str' objects>
```

In Python, class is a synonym for "type" and object is a synonym for "value." The period in str.count tells Python to look for the named method in the class str. 'actg'.count has a similar effect, because the type of 'actg' is str. The statement "every value has a type" can be rephrased as "every object is an instance of a class." Classes and instances form the basis of object-oriented programming. Built-in types and types loaded from Python libraries take advantage of Python's object-oriented features. Chapter 5 discusses how to use those to build your own classes, but until then all we'll need is the little bit of object-oriented vocabulary just introduced.

^{*} Older versions of Python did distinguish classes from types and objects from values.

Assigning Names

A Python program consists of a series of *statements*. Statements do not produce values and cannot be used as expressions.

STATEMENT

Assignment

An assignment statement binds a name to an object. Assignment is denoted by a single equals sign:

```
name = value
```

All an assignment statement does is name a value. That value can also have other names. Assigning a name to a value never changes that value, or any of its other names. Here are some examples of assignment statements and their effects:

```
>>> aaseq1 = 'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'
                                                  # no value printed
>>> aaseq2 = 'AARHQGRGAPCGESFWHWALGADGGHGHAQPPFRSSRLIGAERQPTSDCRQSLQ'
>>> aaseq1
'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'
>>> len(aaseq2)
54
>>> aaseq3
                                                  # never bound
Traceback (most recent call last):
 File "<pyshell#65>", line 1, in <module>
NameError: name 'aaseq3' is not defined
>>> aasea3 = aasea1
>>> aaseq3
'MNKMDLVADVAEKTDLSKAKATEVIDAVFA'
```

Assignment statements may be chained together. This feature is used primarily to bind several names to some initial value:

```
>>> a = b = c = 0
```

Another variation on assignment statements is a bit of shorthand for arithmetic updates. Statements like the following are so common:

```
>>> a = a + 1
```

that Python provides a shorthand for them:

```
>>> a += 1
```

This is called an augmented assignment statement. The operation of an augmented assignment statement can be any of the arithmetic operations shown in the previous chapter: +=, -=, *=, /=, //=, %=, and **=.

Defining Functions

New functions are defined with function definition statements. A definition is a compound statement, meaning it comprises more than one line of code. We will see other kinds of compound statements later, especially in Chapter 4. The first line of each compound statement ends with a colon. Subsequent lines are indented relative to the

The standard practice is to indent by four spaces, and you should adhere to that rule. Also, do not use tab characters since they confuse Python's indentation computations.

STATEMENT

Function Definition

Definition statements have the general form:

def name(parameter-list):

where body is one or more statements, each indented relative to the line with the def. The parameter-list is a (possibly empty) list of names. If there is more than one parameter in the list, they are separated by commas.

When one function calls another, Python records the call from the first function and passes control to the second. If that one calls a third, its call would be recorded and control passed to the third. The record of a call includes the location of the function call in the caller's code so that when a function regains control it can continue executing the code following the function call. Figure 2-3 illustrates how function calling works, with the curved arrows representing the call records.

Some functions are intended to return a value and some aren't. A return statement is used to return a value from a function. The word return is usually followed by an expression whose value is returned as the value of the function. Occasionally a return is not followed by an expression, in which case the value returned is None. All functions return a value, whether or not they contain a return statement: if the function finishes executing without encountering a return, it returns None.

STATEMENT

Function Return

return exits the currently executing function, returning value to its caller. value may be omitted, in which case None is returned.

return value

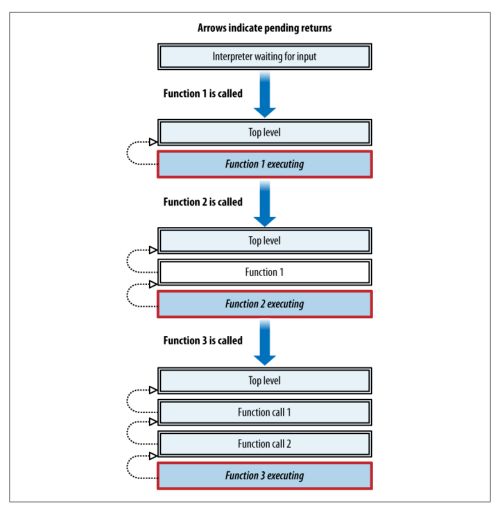


Figure 2-3. Function calls



If you call a function from the interpreter or an IDE while you are developing code and nothing is printed, that means your function returned None. A very common reason for this is that you simply forgot to include a return statement in the function definition. Even experienced Python programmers stumble over this from time to time.

When a function returns, the function that called it continues executing at the point in its code following the call. This happens whether the function returns by executing all its statements or by executing a return statement. Once the function returns, Python no longer needs the information associated with its call. Figure 2-4 illustrates the returns from the functions shown in Figure 2-3.

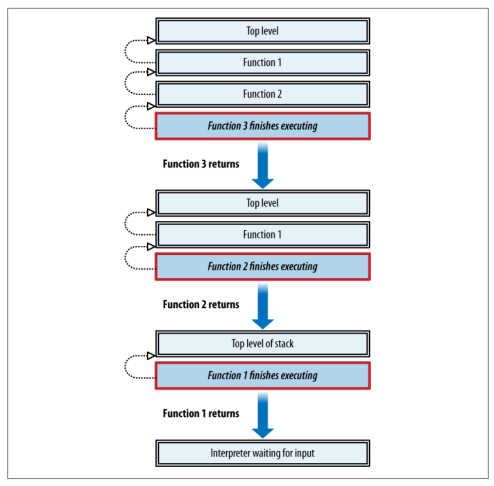
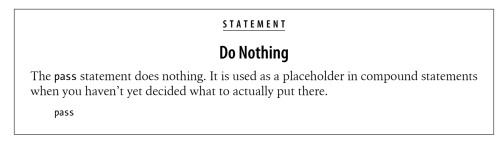


Figure 2-4. Function returns

When you start working on a function, you might not know what to put in the body until you've done some more experimentation. A function's body must contain at least one statement. The do-nothing statement that is equivalent to the no-value None is the simple statement pass.



A minimal function definition would therefore be:

```
def fn():
    pass
```

Now fn can be called, though it does nothing and returns nothing:

```
>>>
```



Words such as def, return, and pass are known as keywords or, because programs cannot use them as names, reserved words. True, False, and None are keywords, as are operators such as and, or, not, and in. A complete list of keywords is found in Table A-1 in Appendix A.

Function Parameters

When a function is called, Python evaluates each of the call's argument expressions. Then it assigns the first parameter name to the first argument's value, the second parameter name to the second value, and so on, for however many parameters there are. Once all the parameter names have been assigned, the statements in the function body are executed. Example 2-1 shows a definition based on a simple use of str.find.

Example 2-1. A simple function for recognizing a binding site

```
def recognition site(base seq, recognition seq):
   return base seq.find(recognition seq)
```

Each function has a separate namespace associated with it. The function's parameters go in that namespace. Assignment statements in the function body also create bindings within the function's namespace. Therefore, a name bound in the interpreter and a name with the same spelling bound in a function definition are in different namespaces and have nothing to do with each other. Example 2-2 illustrates these details.

Example 2-2. Some function definition details

```
def validate base sequence(base sequence): 1
   seq = base sequence.upper() @
   return len(seq) == (seq.count('T') + seq.count('C') + 3
                        seq.count('A') + seq.count('G')) 4
>>> seq = 'AAAT'
>>> validate base sequence('tattattat')
True
>>> seq
'AAAT'
```

Line **1** begins the definition of a function called validate base sequence that has one parameter, base sequence. Line 2 assigns the name seq to the result of calling the method upper on the value of base sequence. The purpose of line 2 is to simplify the rest of the definition by ensuring that all the base characters will be uppercase even if the function is called with a lowercase string. By assigning a name to the result of upper, we avoid having to repeat that call four times later.

Lines 3 and 4 compare the length of the sequence string to the sum of the number of Ts, Cs, As, and Gs in the sequence string. If the length is equal to that sum, the function returns True; otherwise, there is a character in the parameter sequence that isn't a valid base character, and the result will be False.

Note that the right side of the equality expression beginning on line 3 continues on line 4. This is because it begins with a left parenthesis, and Python won't consider it complete until it sees a matching right parenthesis. Without the opening parenthesis, the interpreter would produce an error message as soon as it encountered the end of line 2. Line breaks can appear within a parenthesized expression, giving you more control over how you format long expressions.

Another way to continue a long line without using parentheses is to end the line with a backslash; this tells Python that the current line is continued on the next. (Usually it's preferable to rely on parentheses for multiline expressions, but occasionally backslashes are a better choice.) Example 2-3 shows Example 2-2 rewritten using a backslash instead of the outermost parentheses.

Example 2-3. Using backslash for line continuations

```
def validate base sequence(base sequence):
   seq = base sequence.upper()
   return len(seq) == \
                seq.count('A') + seq.count('G') + \
                seq.count('T') + seq.count('C')
```

The definition is followed by a blank line. What is significant is not that the line is blank but that it is not indented to a position beyond that of the def that begins the definition. A block in Python ends with the first line that is indented less than the first line of the block—i.e., the line following the one that ends with a colon.

Comments and Documentation

The # character tells Python to ignore the rest of the line. This convention can be used for a short comment following the code on a line, or for a more extensive comment consisting of one or more lines beginning with a #. For instance, we could add comments to the validate_base_sequence definition, as in Example 2-4.

Example 2-4. Documentation of a function with commented lines

```
# A short example illustrating various details of function definitions. Given a string ostensibly
# representing a base sequence, return True or False according to whether the string is composed
# entirely of upper- or lowercase T, C, A, and G characters
def validate base sequence(base sequence):
    # argument should be a string
    seq = base sequence.upper()
    # ensure all uppercase characters
```

```
return len(seq) == (seq.count('T') + seq.count('C') +
                    seq.count('A') + seq.count('G'))
```

In general, the kind of information provided in the comment lines before the function definition in this example should be inside the definition, not in a comment before it. Python has a separate feature for this called a *docstring*. Syntactically, it is simply a string that begins the function definition. (Because it is often more than one line long, triple quotes—single or double—are normally used.) Example 2-5 shows the comment of Example 2-4 changed to a docstring.

Example 2-5. A function definition with a docstring

```
def validate base sequence(base sequence):
    """Return True if the string base_sequence contains only
    upper- or lowercase T, C, A, and G characters, otherwise False"""
    seq = base seq.upper()
    return len(seq) == (seq.count('T') + seq.count('C') +
                          seq.count('A') + seq.count('G'))
```

Normally, the only kinds of expressions that have any effect when used as statements are function and method calls; docstrings are an exception. Docstrings are different from comments. Comments disappear when Python interprets code, but docstrings are retained. Consequently, docstrings have greater utility than comments. In particular, the help function looks at the docstring of a user-defined function together with its parameter list to generate a help description, as shown in Example 2-6. (The phrase in module main simply means "defined in the interpreter.")

Example 2-6. Help called on a user-defined function

```
>>> help(validate base sequence)
Help on function validate base sequence in module main :
validate base sequence(base sequence)
   Return True if the string base sequence contains only
   upper- or lowercase T, C, A, and G characters, otherwise False
```



Function definitions contain only statements. A string by itself is a statement because it is a value, a value is an expression, and an expression is a statement. However, docstrings are the only place where a freestanding string would have any effect.

Let's look at another example. Genomes of different species—and regions within a genome—vary with respect to the proportion of Gs and Cs in their DNA as opposed to Ts and As. It is a straightforward exercise to compute the GC content of a given DNA sequence represented as a string. One of the advantages of an interactive environment such as Python's interpreter is that you can experiment with small pieces of code before putting them together. Here's what such experimentation might look like in working toward a definition of a gc content function:

```
>>> seq = 'ATCCGGGG'
>>> seq.count('G')
>>> seq.count('C')
>>> len(seq)
>>> seq.count('G') + seq.count('C')
>>> (seq.count('G') + seq.count('C')) / len(seq)
```

The end result is shown in Example 2-7.

Example 2-7. Defining a function to compute GC content

```
def gc content(base seq):
    """Return the percentage of G and C characters in base_seg"""
    seq = base seq.upper()
    return (seq.count('G') + seq.count('C')) / len(seq)
```

A meaningful example of the use of this function would require a long base sequence, and then it would be difficult to tell whether the result was correct. When writing functions, it is generally a good idea to construct some small "test cases" for which the correct answer is known:

```
>>> seq50 = 'AACCTTGG'
>>> seq75 = 'ATCCCGGG'
>>> seq40 = 'ATATTTCGCG'
>>> gc content(seq50)
0.5
>>> gc_content(seq75)
0.75
>>> gc content(seq40)
0.4
```

Assertions

While developing code, it often happens that your functions get called with arguments of the wrong type or value. Depending on the details of the code, an invalid argument value may result in a Python error or a nonsensical result. Even worse, it might result in an apparently meaningful, but incorrect, result. For example, the definition of gc content assumes that it gets called with a string that contains only (upper- or lowercase) Ts, Cs, As, and Gs. Since it is only counting Cs and Gs, all other characters are assumed to be Ts or As. If given a string that contains a few invalid characters, it will return a value that "makes sense" but is nonetheless "off" by a small amount.

It's often a good idea to express a function's assumptions in its docstring. However, the documentation doesn't have any effect on the computation—nothing stops someone from calling a function with values that violate the documented assumptions. To ensure compliance Python provides a simple assertion statement, one of whose uses is to validate arguments.

STATEMENT

Assertion

An assertion statement tests whether an expression is true or false, causing an error if it is false.

assert expression

Example 2-8 illustrates what happens when an assertion fails.

Example 2-8. A failed assertion

```
>>> assert 1 == 2
Traceback (most recent call last):
 File "<pyshell#53>", line 1, in <module>
    assert 1 == 2
AssertionError
```

There is another form of the assertion statement that takes two expressions. If the first expression is false, the second becomes part of the error report.

STATEMENT

Two-Expression Assertion

A two-expression assertion statement takes two arguments: an expression to evaluate and an expression to use in the error report if the first expression is false.

assert expression1, expression2

Example 2-9 shows a two-expression assertion.

Example 2-9. A failed assertion with a second expression

```
>>> assert 1 == 2, 'invalid arguments'
Traceback (most recent call last):
  File "<pyshell#54>", line 1, in <module>
    assert 1 == 2, "invalid arguments"
AssertionError: invalid argument
```

Let's improve gc content by ensuring that its argument is a valid sequence string. We already have a function that does what we need—validate base sequence. We'll add an assertion statement that calls that function. The assertion in Example 2-10 both documents the function's assumptions and enforces them when the function is called.

```
def gc content(base seq):
     ""Return the percentage of G and C characters in base_seq"""
   assert validate base sequence(base seq), \
           'argument has invalid characters'
   seq = base seq.upper()
   return ((base seq.count('G') + base seq.count('C')) /
            len(base seq))
```

The function in Example 2-10 calls another function that we've defined. This is quite common. Each function should perform what is conceptually a single coherent action. This means you'll write many relatively small functions that call each other rather than writing large functions with many steps.

You can think of the functions you define as building up a vocabulary. In defining a vocabulary, it is natural to define some of its words in terms of others. Python provides a small initial set of words, and you expand on that set by defining more of them.

One reason for writing small functions is that simple functions are easier to write and test than complicated ones. If function A calls B, and B calls C, and C calls D, but something isn't working in D, you can call D yourself from the interpreter. Once you're sure that D works, you can call C, and so on. Another reason for defining a separate function for each action is that a very focused function is likely to prove more useful in other definitions than a larger multipurpose one.

Writing many small functions helps avoid code duplication, that is, having a group of statements repeated in more than one definition. Code duplication is bad because if you want to change those statements you'll have to find every definition that uses them and spend considerable time making the same changes in each definition. Changing multiple occurrences risks ending up with inconsistencies.

Default Parameter Values

You'll often find that most calls to a certain function are likely to include the same value for a particular parameter. Frequently this value is something simple such as True, 0, or None. In such cases, Python provides a way to assign a default value to the parameter that will be used if no explicit value is included in a call to the function.

As mentioned in the preceding chapter, some parameters of some functions are optional. A function definition designates an optional parameter by assigning it a default value in the parameter list. There can be any number of parameters with default values. All parameters with default values must follow any parameters that do not have them.

By way of example, we'll make validate base sequence more flexible by giving it the ability to handle RNA sequences too. This simply entails checking for a U instead of a T. Example 2-11 adds a second parameter whose value determines whether the function looks for Ts or Us. The term or syllable flag (sometimes abbreviated to flg) is commonly used for a Boolean value that determines which of two actions is performed. We'll call our new parameter RNAflag. By using a three-part conditional expression, we can make the change with very little rewriting.

Example 2-11. Adding a "flag" parameter

```
def validate base sequence(base sequence, RNAflag):
    """Return True if the string base_sequence contains only upper- or lowercase
    T (or U, if RNAflag), C, A, and G characters, otherwise False"""
    seq = base sequence.upper()
    return len(seq) == (seq.count('U' if RNAflag else 'T') +
                         seq.count('C') +
                         seq.count('A') +
                         seq.count('G'))
>>> validate_base_sequence('ATCG', False)
>>> validate base sequence('ATCG', True)
False
>>> validate base sequence('AUCG', True)
True
```

The new definition requires calls to the function to provide two arguments. To make using the function a bit more convenient, we can give RNAFlag a default value, as shown in Example 2-12. We don't really know whether the function will get used more often for DNA sequences or RNA sequences, so it's hard to choose a default value on that basis. In general, default values should be chosen to "turn off" the feature a parameter provides, so that users of the function who don't know about the feature aren't surprised by its use. A common default is the kind of false value appropriate for the situation: False, an empty string, a zero, etc. Flags, being Boolean, normally have default values of False.

Example 2-12. Adding a default value for the flag parameter

```
def validate base sequence(base sequence, RNAflag=False):
    """Return True if the string base_sequence contains only upper- or lowercase
    T (or U, if RNAflag), C, A, and G characters, otherwise False"""
    seq = base sequence.upper()
    return len(seq) == (seq.count('U' if RNAflag else 'T') +
                         seq.count('C') +
                         seq.count('A') +
                         seq.count('G'))
>>> validate base sequence('ATCG', False)
>>> validate base sequence('ATCG', True)
False
>>> validate base sequence('AUCG', True)
True
>>> validate base sequence('AUCG')
>>> # second argument omitted; defaults to True
True
```

Keyword parameters have the following properties:

- They are optional (i.e., do not need to be included in calls to the function).
- They are defined by specifying default values in the function's parameter list; if the keyword does not appear in a call to the function, its default value is used.
- In both the parameter list of a function definition and the argument list of its calls, all "positional" (i.e., required) arguments must appear before any keyword parameters.
- Keyword parameters may appear in any order.
- With a few exceptions, keyword arguments may be provided positionally in a function call in the order in which the keywords appear in the function's definition.

Using Modules

In addition to those built into the base interpreter environment, Python offers a large selection of optional types, functions, and methods. These are defined by *module* files placed in a library directory as part of Python's installation. Modules can also be obtained from external sources and added to Python's library.

A module file is an ordinary file containing Python statements (defs, mostly). A docstring at the beginning of the file (if present) describes its contents and purpose. Later chapters of this book discuss many modules important for bioinformatics programming. Here, I'll only explain the mechanics of module use and give a few basic examples.

Importing

A module's contents are brought into the interpreter's environment by an import statement.

STATEMENT

Importing

The basic form of the import statement loads a module into the Python environment and makes its name available in the namespace into which it was imported (usually the interpreter or a program).

import name

The *name* is just the name of the module—no path and no extension.

For example, the module os provides an interface to the computer's operating system. You can import it using this statement:

>>> import os

Python keeps track of the directories in which modules may be found and of which modules have already been imported. The first time a module is imported—with any

form of import statement—Python looks for it in each library directory in turn until it is found. It then executes the file's statements and creates an object to represent the module and its contents. Subsequent imports skip all of this.

Python incorporates quite a few modules during its build process. While their contents are in the system, their names are not. Consider the os module just mentioned all the contents of the module are actually already in Python, but the name os has no meaning until the module is imported. Importing assigns the name of the module to the object that represents the module's information.

Module namespaces

Each module has a separate namespace associated with it. In fact, a module object is little more than a namespace. Module contents are accessed using the same dot notation used for method calls. Some modules even have submodules, which are referenced using dot notation—for example, os.path. Let's look at some examples:

```
# after import
<module 'os' from '/usr/lib/python31/os.py'>
>>> os.getcwd()
                                                     # a function in the os module
'/Users/mlm/programming/python'
>>> os.getlogin()
                                                     # another function in the os module
'mlm'
```

Each module's namespace is isolated from the interpreter's namespace and the namespaces of other modules. Functions with the same name can be defined in multiple namespaces and not interfere with each other, because each is accessed by a name in a different namespace.

Another form of the import statement is used to purposely bring a name from a module into the namespace in which the import appears—the interpreter or another module.

STATEMENT

Selective Import

This form of import statement loads a library into the Python environment but does not make its name available in the namespace from which it was loaded. Instead, it imports specific names from the module into the importing namespace.

```
from modulename import name1, name2, ...
```

You can give something you are importing a different name when you import it by using the following form of the import statement:

```
from modulename import actualname as yourname
```

You can import *all* the names from a module with the following variation:

```
from modulename import *
```



In general, it is best to avoid the "import all" form (occasionally a module's documentation or examples suggest you begin a program by importing all of its names, in which case it's probably all right to do so). The reason to avoid this form is that you don't know what names you'll be importing. Some might rebind existing names—this is called a *name conflict*—and lead to strange problems. Even importing specific names using the from form isn't necessarily the best idea—when someone else reads your code, or you read it later, using the module prefix makes it clear which names come from which modules and, if there is no prefix,

If your code frequently uses a particular name from a module, you'll find it convenient to import that name into the code's namespace. For example, sys is another module that is already in the base Python system without its name being in the interpreter's namespace. One name sys provides is version, whose value is a string describing Python's version. After executing the following statement:

```
from sys import version
```

your code will be able to refer directly to version instead of sys.version. Consider the interactions in Example 2-13.

Example 2-13. Importing and namespaces

are defined in the file itself.

```
① >>> svs
Traceback (most recent call last):
  File "<pyshell#103>", line 1, in <module>
NameError: name 'sys' is not defined
2 >>> version
Traceback (most recent call last):
  File "<pyshell#105>", line 1, in <module>
NameError: name 'version' is not defined
3 >>> from sys import version
4 >>> version
'3.0 (r30:67503, Jan 2 2009, 12:13:58) \n[GCC 4.0.1 (Apple Inc. build 5488)]'
6 >>> sys
Traceback (most recent call last):
  File "<pyshell#108>", line 1, in <module>
NameError: name 'sys' is not defined
6 >>> import sys
7 >>> from sys import version number
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
ImportError: cannot import name version number
3 >>> import version
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
ImportError: No module named version
```

This series of inputs demonstrates the following points:

- **1** The name sys is not in the interpreter's namespace.
- 2 The name version is not in the interpreter's namespace.
- **3** The name version is imported from the module sys.
- **1** The name version is now in the interpreter's namespace.
- **5** The name sys is still not in the interpreter's namespace.
- **6** The name sys is now in the interpreter's namespace.
- The module sys does not define anything called version number.
- **3** There is no module named version.

The random module

One useful module is random, which provides various ways to generate random numbers. The function random.randint takes two integer arguments and returns a random integer in the range of the first to the second inclusive. For example, random.randint(1,4) will return 1, 2, 3, or 4. (Note that the range is inclusive of the second number, unlike with string slicing, which designates a substring from the first position up to but not including the second position indicated by the slice.)

One good use of random.randint is to generate examples and "test cases" to use with code you are developing. Example 2-14 shows how to generate a random codon.

```
Example 2-14. Generating a random codon
```

```
from random import randint
def random base(RNAflag = False):
   return ('UCAG' if RNAflag else 'TCAG')[randint(0,3)]
def random codon(RNAflag = False):
   return random base(RNAflag) + random base(RNAflag) + random base(RNAflag)
```

Notice how breaking the code into two separate functions, each of which does its own narrowly focused job, helps readability. There's not much reason to avoid using small functions like these. You'll see as you write them that they'll turn out to be useful for other code. It's helpful to think of types, functions, and methods as parts of a vocabulary: types and their instances are nouns, and functions and methods are verbs. Your definitions add to that vocabulary.

Example 2-15 shows a function that simulates single-base mutation. It uses assignment statements to name everything along the way to the final result.

```
from random import randint
def replace base randomly using names(base seq):
    """Return a sequence with the base at a randomly selected position of base_seq replaced
    by a base chosen randomly from the three bases that are not at that position""
    position = randint(0, len(base seq) - 1)
                                                     # -1 because len is one past end
    base = base seq[position]
    bases = 'TCAG'
    bases.replace(base, '')
                                                      # replace with empty string!
    newbase = bases[randint(0,2)]
    beginning = base seq[0:position]
                                                      # up to position
```

Example 2-16 shows another version of the same function. This variation uses a complex expression instead of a lot of names. It does use the name position to save the value of the call to randint, because that is used in two places; if randint were called twice, it would return two different values, and the definition needs to use the same value in both places.

omitting the base at **position**

Example 2-16. The same function without intermediate result names

end = base_seq[position+1:]

return beginning + newbase + end

```
def replace base randomly using expression(base seq):
   position = randint(0, len(base seq) - 1)
   return (base seq[0:position] +
            'TCAG'.replace(base seq[position], '')[randint(0,2)] +
           base seq[position+1:])
```

There's no clear-cut rule about which style to use. Sometimes naming things improves readability (even taking the place of comments) and aids debugging. Often, however, it doesn't contribute much, especially if the value named is only used once in the definition. In the case of this function, something between these two extremes would probably be best. The compromise version is shown in Example 2-17.

Example 2-17. A function definition with some intermediate names

```
def replace base randomly(base seq):
   position = randint(0, len(base seq) - 1)
   bases = 'TCAG'.replace(base seq[position], '')
   return (base seq[0:position] +
           bases [randint(0,2)] +
           base seq[position+1:])
```

Python Files

A file of yours that you import becomes a module just as if the file had been in Python's library. Accessing any of its contents requires using the file's name as a prefix, except for specific names imported into the interpreter's namespace using the "from" form of the import statement. Normally a file meant for importing assigns names and defines

functions but doesn't do any observable computation when imported. Files intended to be run or executed normally end with function calls and statements that do something observable, such as printing results.

Each Python file must import any modules it uses. That is, if file A uses module M, it must import module M even if it imports another file that imports M or is imported into the interactive interpreter after M has been imported there. Example 2-18 shows a file containing the functions defined in examples earlier in this chapter, along with a new one for testing the others. The test function is called at the end of the file. That call occurs regardless of whether the file is imported or executed.

Example 2-18. A Python file

```
def validate base sequence(base sequence, RNAflag = False):
    ""Return True if the string base sequence contains only upper- or lowercase
    T (or U, if RNAflag), C, A, and G characters, otherwise False"""
    seq = base sequence.upper()
    return len(seq) == (seq.count('U' if RNAflag else 'T') +
                         seq.count('C') +
                         seq.count('A') +
                         seq.count('G'))
def gc content(base seq):
     ""Return the percentage of bases in base_seq that are C or G"""
    assert validate base sequence(base seq), \
            'argument has invalid characters'
    seq = base seq.upper()
    return (base_seq.count('G') +
            base seq.count('C')) / len(base seq)
def recognition site(base seq, recognition seq):
   """Return the first position in base_seq where recognition_seq
   occurs, or -1 if not found"""
   return base seq.find(recognition seq)
def test():
    assert validate base sequence('ACTG')
    assert validate base sequence('')
    assert not validate base sequence('ACUG')
    assert validate base sequence('ACUG', False)
    assert not validate base sequence('ACUG', True)
    assert validate base sequence('ACTG', True)
    assert .5 == gc content('ACTG')
    assert 1.0 == gc content('CCGG')
    assert .25 == gc content('ACTT')
    print('All tests passed.')
test()
```

Tips, Traps, and Tracebacks

Tips

Names

- One object may have many names.
- Choose clear, descriptive names for functions and arguments.
- Assignment changes what a name means—i.e., which object it references. If a name refers to something that is modifiable (a phenomenon we will encounter in the next chapter), assignment does not modify the object.

Function definitions

- Think of function definitions as adding to Python's vocabulary of built-in functions, types, methods, etc.
- Use pass as a placeholder for the body of a function you are defining; it allows you to start the function definition and specify its parameters without it doing anything while you work on the rest of the code. The other code can even call the function at the appropriate place. Later, you can replace the pass to define what the function should really do.
- Your function definitions should normally be quite small. Each should do one main thing. Get comfortable with the idea of defining functions that call each other.
- Except perhaps for functions of a couple of lines with self-descriptive names, always include docstrings in your function definitions.
- Read the following (and occasionally reread them, as they will make more sense and seem more helpful the more Python work you do):
 - —Style Guide for Python Code
 - —Docstring Conventions
 - The Zen of Python, accessed by typing the following at the command line:

import this



The code examples in this book do not adhere to certain aspects of the official docstring conventions. In particular, the conventions call for docstrings longer than one line to begin with a one-line summary and a blank line before continuing with further details, and to end with a blank line followed by the closing quotes on a separate line. Given the large number of examples contained in this book and an approach to writing code that emphasizes very small functions, following the conventions strictly would have resulted in docstrings whose size rivaled that of the code they documented, as well as simply taking up too much space on the page.

Executing code

- Do not be intimidated by tracebacks—they identify the nature and location of the problems they are reporting.
- Start putting your code in Python files as soon as possible. Testing bits of code in the interpreter and then copying them into a file is fine, but eventually, as you become more competent in writing Python code, you will want to edit files directly and then run them in the interpreter. You can use whatever text editor you are comfortable with. Save the files in text format with the extension .py.
- Learn how to run Python programs from the command line. If you don't know how, ask someone to show you. (They don't have to know Python!) Using the file *echo.py* as an example, and assuming that the python3 command runs Python:
 - 1. Type:

```
pythonpath/python3 echo.py
```

where *pythonpath* is the full path to the Python executable on your machine.

2. If the folder containing the Python executable is in your execution path, as defined by system and shell variables, you only need to type:

```
python3 echo.py
```

These commands will execute the code in echo.pv, then exit. To execute all the code in the file, enter the interpreter and include -i (for "interactive") before the name of your file.

- One of the great advantages of using an interactive interpreter is that you can test pieces of your code without having to test entire function definitions or scripts:
 - —When you are first defining a function, you can try out bits of code or aspects of Python about which you are uncertain.
 - —You can try various ways of expressing a part of a function definition.
 - —After assigning names, you can look at their values.
 - —You can test functions interactively by calling them with various values.

Using IDLE

The Python installation comes with a simple integrated development environment (IDE) called IDLE. It has a window with the interpreter running in it and it allows you to edit files in other windows. You can start it by double-clicking its icon or by doubleclicking a Python file. (You may have to change the default application for .py files to open with the Python 3 version of IDLE: on Windows, the default is to run the program in a shell, then exit the shell. On any platform, if a version of Python 2 is present on your computer, you may have to change the default application for .py files to be the Python 3 version of IDLE.

When you open a file with IDLE—either by double-clicking it or selecting the Open command from IDLE's File menu—the file will be displayed in an IDLE editor window. Editor windows provide many conveniences, including "syntax highlighting" and keyboard shortcuts. However you open IDLE, you will see a "Python Shell" window that is the equivalent of the command-line interpreter. Here are some hints for using IDLE:

- The typical shortcut keys are available for the Edit menu's commands, for most of the other menu commands, and for some movement and editing commands that are not on a menu. The Preferences command on the IDLE menu opens a tabbed window that you can use to customize IDLE's appearance (including the typeface and font size), shortcut keys, and behavior.
- The IDLE Help command on the Help menu brings up a window explaining the menu commands, keyboard shortcuts for editing and moving around, and other aspects of using IDLE. (This is also available as the file help.txt in the idle subdirectory of the Python installation's library directory.) † You should periodically review this documentation, as IDLE has many useful features that you won't at first appreciate.
- When in an editor window, you can execute the code in the window by selecting the Run Module command from the Run menu or pressing F5. (The Run menu is replaced by the Debug menu when the cursor is in the interpreter window.) Every time you execute code using IDLE's Run Module command, the Python interpreter is restarted. No previous assignments, definitions, or imports survive this restart. The restart behavior is another reason why it's useful to include everything you need in files you are developing in IDLE, including assignments to strings and numbers that you want to use in testing and debugging your code. Basically, anything you don't want to type repeatedly should go in a file. However, once you've run the code, all of its assignments and definitions will be available in the interpreter until the next time you execute the Run Module command.
- The text in IDLE's editor window is colored to distinguish various aspects of Python code. (The colors are also customizable, so if you find some either too dramatic or too difficult to distinguish, you should take the time to adjust them to your liking.) This feature is called, naturally enough, syntax highlighting. It makes the code much more readable. In addition, the color of the text you are presently typing and of the text that follows it can reveal syntax problems without you even running the program.

Perhaps the most useful example of this is when you notice that the text you are typing is colored as a string, indicating that you did not type the quotes required to end the string where you intended. Incomplete multiline strings (those beginning with three single or three double quotes, as with docstrings) make for especially dramatic effects: as soon as you begin the string, the rest of the file gets recolored

[†] See also section 24.6 of the Python library documentation (library/idle/idle.html in the HTML version).

as a string—at least, until the next set of matching triple quotes is encountered. That's fine if you're still typing the string, but if the following code is still colored as a string after you think you have finished typing the string, you need to add or fix the quotes at the end of the string.

- A particularly helpful command is Show Surrounding Parens on the Edit menu, typically used by typing its keyboard shortcut. This briefly highlights the code around the cursor within the corresonding parentheses, brackets, or braces. This can help you spot mistakes and clear up confusion about the structure of expressions you are editing.
- When you press Return (Enter), IDLE will indent the new line to a column matching the current syntactic context. For example, when you hit Return after the colon at the end of the first line of a function definition, it indents the next line four spaces. If the line doesn't start in the column you expect, check the syntax of the previous lines. Common situations in which this occurs include:
 - —When you type more right parentheses at the end of a statement than are needed to "close" the last expression of the statement. The next line will indent much further to the *left* than you expected, perhaps to column 0.
 - —When you don't type enough right parentheses to close the last expression. The line will indent much further to the *right* than you expected.
- Another very useful feature of IDEs is automatic completion of partially typed Python names and filenames. Taking advantage of automatic completion can save you not just a lot of typing, but a lot of mistakes (down to either typos or misremembering Python names). Here are some autocompletion hints (be aware that it can take a while to get used to how this facility works):
 - —If you pause for a few seconds after typing some characters, and there are possible completions to what you have just typed, an "autocomplete window" opens.
 - —Typing a tab brings up the autocomplete window immediately.
 - —If there is only one possibility, the word will be completed without the window appearing.
 - —The IDLE Preferences window allows you to customize several aspects of autocompletion.

Eventually, if you end up doing heavy-duty software development, you might want to switch to a more sophisticated environment. There are a number of them to choose from, including individual efforts, open source projects, and free, "personal," and "professional" editions of commercial products. As of this writing, only a small number have been released in a version compatible with Python 3. This is another reason to use IDLE—it is part of Python itself, so the version that comes with the Python 3 installation works with Python 3.



You can find a complete list of Python IDEs at http://wiki.python.org/ moin/IntegratedDevelopmentEnvironments. There is no indication in that list of which IDEs support Python 3, but those last updated before 2009 definitely do not (unless the page's information is not current).

Managing Python files

- Filenames are expected to be in all lowercase and to have the extension .py.
- Python files may be edited in any text editor as long as they are saved in plain text
- Do *not* save the files you edit in Python's or IDLE's directory, for these reasons:
 - Other people may be using the Python installation on your machine; seeing your files there may confuse them, and they may even delete or overwrite them.
 - You may want to replace your entire Python installation; in this event, you won't want to have to figure out which files are yours and move them someplace else before you can do so.
 - —Eventually you will want to have multiple directories for your Python files different projects, different versions of a project, etc.
 - Instead, create a Python working directory wherever you feel comfortable putting it—in your *Documents* folder, for example. Then create a subdirectory for each project.
- Back up the directory containing all your Python work daily, if not more often. As you work, save your files frequently according to a set backup scheme; for instance, if working on test.py, save it to test01.py, then a bit later to test02.py, etc.[‡] (The zero is so that when you get to version 10, file listings will still be alphabetized.) Saving multiple versions will make you less wary of experimenting, since you'll know you can always go back to an earlier version—if you decide to abandon the direction you took while working on part of your code, it is often easier to revert to an earlier version than it is to make wholesale changes to the current one.
- Files with the extension .pyc are intermediate files that contain "compiled" versions of the corresponding .py file's definitions. Python saves these after successfully importing a .py file for the first time, and after importing a file that has been changed since the last time it was imported. The only purpose of these files is to increase import speed—they have no effect on how fast programs actually run once their modules have all been loaded.

[‡] If you happen to know how, or can get someone to show you, a better approach is simply to manage the files with a version management system such as CVS, Subversion, git, or Mercurial.



It is always safe to delete the .pyc files. In fact, deleting old .pyc files can be a good idea. You may occasionally rename or delete .py files and then accidentally import a module with one of the old names. The import will be successful as long as there is still a corresponding .pyc file—that is, your code might import a module you thought was gone. Of course, you won't have meant for your code to import a deleted or renamed module: the issue here is that it can happen in a way that doesn't produce an error message to alert you to the need to change the import statement.

Traps

- Assignment uses one =, comparison two (==). Using only one instead of two is a very common error. Worse, it often occurs in a context where it isn't a syntax error, in which case you are left to figure out the program's consequent misbehavior by vourself.
- If your function doesn't do anything when you call it, you probably forgot to include a return statement in it.
- If nothing happens when you run a file, either from the command line or within IDLE, your file contains no top-level statements that produce output. The file should end with calls to print or functions you've defined that print things.
- Names assigned in a function have no meaning outside that function. Two different functions can assign the same name, but because the names are in different scopes they have no relation to each other.
- Keyword arguments must appear after all the "positional" (i.e., required) arguments in a function call.
- Module names are not quoted in import statements, nor do their names include .py.
- Syntax errors are frequently caused by something wrong at the end of the line before the line shown by the error message. Common causes of these errors include:
 - —Failing to end the first line of a def with a colon
 - —Ending the previous line with too few or too many parentheses

Before even thinking about what the error means, you should glance at the line indicated by the error message, and then the one before it. If you notice an obvious problem, fix it and try again.

Tracebacks

Here are some representative error messages:

AssertionError: invalid argument

The value in an assertion statement is false; the second value in the statement, if there is one, becomes the message.

ImportError: No module named version

An attempt was made to import a module named version, but Python couldn't find a module by that name.

ImportError: cannot import name vision

An attempt was made to import a name from a module using the from module import name form of the import statement, but the module doesn't define anything with that name.

NameError: name 'Non' is not defined

Python doesn't recognize a name: it's not built in and hasn't been defined with an assignment or def statement. Other than misspellings, two frequent causes are the use of the name in one function when it was assigned in another and the use of a name from a module without preceding it with the module's name or importing the name from the module.

TypeError: gc content() takes exactly 1 positional argument (0 given)

A call to a function supplied too few or too many positional (i.e., required, or, equivalently, nonkeyword) arguments.

Collections

Instances of primitive Python types represent individual values, as we saw in the first chapter. Python also has built-in types that are *compound*, meaning that they group together multiple objects. These are called *collections* or *containers*. The objects in a collection are generally referred to as *items* or *elements*. Collections can hold any number of items. Some collection types can even contain items with a mixture of types, including other collections.

There are three categories of built-in collection types in Python: *sets*, *sequences*, and *mappings*. *Streams* are another kind of collection whose elements form a series in time, rather than the computer's memory space. Though they are not formally recognized in Python, many of Python's functions and methods—especially those that manipulate files—implement stream-like behavior. Modules such as array and collections provide additional collection types.

The primary distinguishing characteristic of the four collection categories is how individual elements are accessed: sets don't allow individual access; sequences use numerical *indexes*; mappings use *keys*; and streams just provide one address—"next." Strings straddle the border between primitives and collections, having characteristics of both; when treated as collections they are interpreted as sequences of one-character substrings.

Some types are *immutable*, meaning that once they are created they cannot be changed: elements may not be added, removed, or reordered. Primitive (noncollection) types are also immutable; in particular, strings cannot be changed once created. There is one very important restriction on the types of elements some collections can contain: mutable built-in collections cannot be added to sets or used as mapping keys.

You can create a collection by calling its type as a function. With no arguments, an empty collection is created. With a sequence argument, a new collection is created from the sequence's elements. Some collection types can be called with an integer argument to create an empty collection of that number of elements. Many of the collection types can also be created directly with Python syntax, the way strings are created using quotes.

Certain basic operations and functions are applicable to all collection types. These are shown in Table 3-1. The first four return a Boolean value; such functions are called predicates.

Table 3-1. Operations and functions that work for all collection types

Operation	Returns
x in coll	True if <i>coll</i> contains <i>x</i> .
x not in coll	True if <i>coll</i> does not contain <i>x</i> .
any(<i>coll</i>)	True if any item in <i>coll</i> is true, otherwise False.
all(coll)	True if every item in <i>col1</i> is true, otherwise False.
len(coll)	The number of items in $coll$ (not supported by streams).
<pre>max(coll[, key=function])</pre>	The maximum item in coll, which may not be empty. ^a
<pre>min(coll[, key=function])</pre>	The minimum item in coll, which may not be empty. ^a
sorted(coll[, keyfn] [, reverseflag])	A list containing the elements of <i>coll</i> , sorted by comparing elements or, if <i>keyfn</i> is included in the call, comparing the results of calling <i>keyfn</i> for each element; if <i>reverseflag</i> is true, the ordering is reversed. <i>keyfn</i> and <i>reverseflg</i> must be specified as keyword arguments, not positionally. ^a

a Atypically, the optional arguments to max, min, and sorted may not be supplied positionally in a call—they must be provided as keyword arguments. The keyfn parameter is explained at the end of the chapter, in the section "Functional Parameters" on page 89.

Sets

A set is an unordered collection of items that contains no duplicates. The implementation enforces the restriction against duplication: attempting to add an element that a set already contains has no effect. The type frozenset is an immutable version of set. There is one important limitation on the type of objects that sets and frozensets can contain: they cannot contain instances of mutable types, or even immutable types that contain mutable elements.

Since strings behave as collections, a string can be used as the argument for a call to set. The resulting set will contain a single-character string for each unique character that appears in the argument. (This behavior may surprise you from time to time!) The order in which the elements of a set are printed will not necessarily bear any relation to the order in which they were added:

Sets can also be created syntactically by enclosing comma-separated values in curly braces ({}). This construction is called a *set display*. There is no corresponding syntax for frozensets—you must call frozenset to create one. Empty braces do not create an empty set, but calling set with no arguments does. (Empty braces create an empty dictionary, discussed shortly.)



Python 2: The brace notation for creating sets is not supported; you must call set as a function to create a set.

Let's look at some examples:

```
>>> DNABases = {'T', 'C', 'A', 'G'}
>>> DNABases
{'A', 'C', 'T', 'G'}
>>> RNABases = {'U', 'C', 'A', 'G'}
>>> RNABases
{'A', 'C', 'U', 'G'}
```

When a string is inside the pair of braces that creates a set, the string is *not* broken up into its individual characters:

```
>>> {'TCAG'}
{'TCAG'}
>>> {'TCAG', 'UCAG'}
{'UCAG', 'TCAG'}
>>>
```

Sets can be compared as shown in Table 3-2. The operator versions require both operands to be sets, but the method versions allow the argument to be any collection.

Table 3-2. Set comparison operations

Operator	Method	Returns
	<pre>set1.isdisjoint(coll)</pre>	$\label{True} \textbf{True} \textbf{if} \textbf{the set} \textbf{and} \textbf{the argument have} \textbf{no} \textbf{elements} \textbf{in} \textbf{common}$
set1 <= set2	<pre>set1.issubset(coll)</pre>	True if every element of $set1$ is also in $set2$ ($coll$ for the method)
set1 <set2< td=""><td></td><td>True if every element of $set1$ is also in $set2$ ($col1$ for the method) and $set2$ is larger than $set1$ (i.e., $set1$ is a proper subset of $set2$)</td></set2<>		True if every element of $set1$ is also in $set2$ ($col1$ for the method) and $set2$ is larger than $set1$ (i.e., $set1$ is a proper subset of $set2$)
set1>= set2	<pre>set1.issuperset(coll)</pre>	True if every element of $set2$ ($coll$ for the method) is also in $set1$
set1>set2		True if every element of set2 is also in set1 and set1 is larger than set2 (i.e., set2 is a proper subset of set1)

Python's set and frozenset types implement the mathematical operations of set algebra. These use regular operator symbols and, in some cases, equivalent method calls. As with the comparison operations, the operator versions require both operands to be sets, while the methods allow the argument to be any collection. Table 3-3 shows the operators and the corresponding methods.

Table 3-3. Algebraic set operations

Operator	Method	Returns
set1 set2	set1.union(coll,)	A new set with the elements of both set1 and set2 (or set1 and each of the arguments)
set1&set2	<pre>set1.intersection(coll,)</pre>	A new set with the elements that are common to set1 and set2 (or set1 and each of the arguments)
set1 - set2	<pre>set1.difference(coll,)</pre>	A new set with the elements in set1 that are not in set2 (or any of the arguments)
set1^set2	<pre>set1.symmetric_difference(coll)</pre>	A new set with the elements that are in either set1 or set2 (or each of the arguments) but not both

Using sets, we can rewrite validate base sequence more succinctly than with the ponderous conditional expressions used in the last chapter. Example 3-1 illustrates.

Example 3-1. Rewriting validate_base_sequence using a set

```
DNAbases = set('TCAGtcag')
RNAbases = set('UCAGucag')
def validate base sequence(base sequence, RNAflag = False):
    """Return True if the string base_sequence contains only upper- or lowercase
    T (or U, if RNAflag), C, A, and G characters, otherwise False"""
    return set(base sequence) <= (RNAbases if RNAflag else DNAbases)</pre>
```



We could have just written out the set values directly, enclosing eight one-character strings within braces, either in the assignment statements or instead of the assigned names in the function definition. Using set to convert a string meant less typing, though, which is always good. A fundamental fact about programming—or any computer use, for that matter—is that every character you type is a chance to make a mistake!

Not only is this version of the function definition more succinct, but it more directly expresses the function's purpose. The function tests whether all the characters in its argument are in the set of allowable base characters. This definition says that explicitly, whereas the previous definition counted the number of times each base occurred and compared the sum of those counts to the length of the sequence. It worked, but it was a roundabout way of saying something that can be said more simply and clearly.

The algebraic set operations produce new sets. There is also a group of operators and methods for modifying sets, shown in Table 3-4. The operator forms require both operands to be sets, while the methods can take any kind of collection argument.



The assignment statements in Example 3-1 were placed before the function definition, so they only get executed once rather than every time the function is called. It's not likely to matter much in this case, but keep in mind that in general you should avoid repeating computations unnecessarily where convenient. You should never let "efficiency" considerations get in your way while developing code, since you won't know how fast your program runs or which parts are slow—or even whether you'll end up using the part of the code whose efficiency concerns you—until the code is working. That said, it's easy enough to keep simple, obvious things like assignment statements whose values don't change out of loops as a matter of practice.

Table 3-4. Set update operations

Operator	Method	Result
set1 = set2	set1.update(coll)	Updates set1 by adding the elements in set2 (co11 for the method)
set1&=set2	<pre>set1.intersection_update(coll)</pre>	Updates $set1$ to keep only the elements that are in both $set1$ and $set2$ ($col1$ for the method)
set1 -= set2	<pre>set1.difference_update(coll)</pre>	Updates $set1$ to keep only the elements that are in $set1$ but not in $set2$ ($col1$ for the method)
set1 ^= set2	<pre>set1.symmetric_difference_update(coll)</pre>	Updates $set1$ to keep only the elements that are in either $set1$ or $set2$ ($col1$ for the method)
	set1.add(item)	Adds item to set1
	<pre>set1.remove(item)</pre>	Removes item from set1; it is an error if item is not in the set
	<pre>set1.discard(item)</pre>	Removes <i>item</i> from <i>set1</i> if it is present; no error if it is not present

Sequences

Sequences are ordered collections that may contain duplicate elements. Unfortunately, the word "sequence" is frequently used in both Python programming and bioinformatics, not to mention in ordinary writing. Hopefully, the context will make its intended meaning clear.

Because sequences are ordered, their elements can be referenced by position. We've already seen how this works with string indexing and slicing; the same mechanisms work with other sequence types. Table 3-5 summarizes the various cases of slicing expressions. Remember that a negative index counts from just past the last element.

Table 3-5. Summary of sequence slicing

Expression	Returns
seq[i:j]	Elements of seq from i up to, but not including, j
seq[i:]	Elements of seq from i through the end of the sequence
seq[:j]	Elements of seq from the first up to, but not including, j
seq[:-1]	Elements of seq from the first up to, but not including, the last
seq[:]	All the elements of seq —i.e., a copy of seq
seq[i:j:k]	Every $k^{ ext{th}}$ element of seq , from $m{i}$ up to , but not including , $m{j}$

There are six built-in sequence types, as summarized in Table 3-6.

Table 3-6. Sequence types

Туре	Element type	Mutable?	Syntax
str	One-character strings ^a	No	Quotes of various kinds
bytes	8-bit bytes ^b	No	A string with a b prefix
bytearray	8-bit bytes	Yes	None
range	Integers	No	None
tuple	Any	No	Parentheses
list	Any	Yes	Brackets

^a Strings don't really "contain" characters or one-character strings, but for many purposes they behave as if they did.

b Bytes are equivalent to the integers from 0 through 255.



Python 2: Substantial changes were made to the sequence types from Python 2.x to 3.x. The most significant change was that Python 2.x strings contain one-byte "ASCII" characters but Python 3.x strings contain "Unicode" characters. In Python 2 range was a function, but in Python 3 it is a type, replacing what in Python 2 was the type xrange. Several other sequence types were also replaced by new ones with different characteristics.

Table 3-7 lists the operations that are supported by all sequence types, with some clarifying details.

Table 3-7. Generally supported by sequence operations

Operation	Restriction
<, <=, >, >=	Same type operands, except that bytes and bytearray may be compared
==, !=	Any type operands, but always False if the operand types are different
in,not in	
Repetition (*)	

Operation	Restriction
Concatenation (+)	Same type operands, except that a bytes and a bytearray may be concatenated, with the type of the result the type of the lefthand operand
Indexing	
Slicing	

All sequence types except for ranges and files support the methods count and index. In addition, all sequence types support the reversed function, which returns a special object that produces the elements of the sequence in reverse order.

Strings, Bytes, and Bytearrays

Strings are sequences of Unicode characters (though there is no "character" type). Unicode is an international standard* that assigns a unique number to every character of all major languages (including special "languages" such as musical notation and mathematics), as well as special characters such as diacritics and technical symbols. Unicode currently defines more than 100,000 characters. The 128 "ASCII" characters (digits, lower- and uppercase letters, punctuation, etc.) have the same numerical values in both plain ASCII and Unicode representations.

The bytes and bytearray types are sequences of single bytes. They have essentially the same operations and methods as str. One important difference is that with a string, regardless of whether you index an element or specify a slice, you always get back a string. With bytes and bytearrays, although slices return the same type, indexing an element returns an integer from 0 through 255.

Another interesting difference is that when either bytes or bytearray is called with an integer argument, it creates an instance of that length containing all 0s. If called with a sequence of integers with values from 0 through 255, it performs the normal "conversion" kind of instance creation, creating a bytes or bytearray object with those integers as its elements. Note that strings do not contain integers—their components are single character strings—so they can't be the argument to bytes or bytearray.

There are some differences in how each sequence type treats the in and not in operations. When the righthand operand is of type str, the lefthand operator must also be a str, and the operation is interpreted as a substring test. When the righthand operand is of type bytes or bytearray, the lefthand side can be an integer from 0 to 255, or it can be a bytes or a bytearray. If it is an integer, in and not in are membership tests; otherwise, they are subsequence tests, as with str.

The bytes and bytearray types serve many purposes, one of which is to efficiently store and manipulate raw data. For instance, a piece of lab equipment might assign a signal

^{*} See http://www.unicode.org/standard/WhatIsUnicode.html and http://www.unicode.org/standard/principles .html for more information.

strength value of between 0 and 255 for something it is measuring repetitively. The natural way to store these values in Python is to use bytes. Data obtained from outside a program in other ways is often in the form of raw bytes. (We'll see this when we discuss database access, web downloads, and other such topics later in the book.) That is, the bytes meant something where the data came from, but they have no intrinsic meaning inside Python. Interpretation of externally obtained bytes is the program's responsibility.

Strings, bytes, and bytearrays have a very wide range of application in the management and manipulation of data and human-readable text. They provide a much richer repertoire of methods than other sequence types. The more commonly used ones are listed in the following sections. The sections refer only to strings, but bytes and bytearrays support the same methods. Other methods will be discussed a little later, in the section "Sequence-oriented string methods" on page 65.

Creating

We know that a new empty collection can be created by using the name of its type in a call with no arguments. Some collection types can also be called with arguments. Following are the creation calls for str, bytes, and bytearray objects:

str()

Returns an empty string

str(obj)

Returns a printable representation of *obj*, as specified by the definition of the type of obj

bytes()

Returns an empty bytes object

bytes(n)

With *n* an integer, returns a bytes object with *n* zeros

bytes(integer-sequence)

With *integer-sequence* a sequence containing integers between 0 and 255 inclusive, returns a bytes object with the corresponding integers

char(n)

Returns the one-character string corresponding to the integer n in the Unicode system

ord(char)

Returns the Unicode number corresponding to the one-character string *char*

The functions for creating bytearray objects work the same way as the corresponding functions for bytes.

Testing

It is often necessary in processing a string to determine what kinds of characters it "contains." These methods provide convenient ways to determine that for the most common cases:

str1.isalpha()

Returns true if *str1* is not empty and all of its characters are alphabetic

str1.isalnum()

Returns true if *str1* is not empty and all of its characters are alphanumeric

str1.isdigit()

Returns true if *str1* is not empty and all of its characters are digits

str1.numeric()

Returns true if str1 is not empty and all of its characters are numeric, including Unicode number values and names

str1.isdecimal()

Returns true if *str1* is not empty and all of its characters are characters that can be used in forming decimal-radix numbers, including Unicode number values and names

str1.islower()

Returns true if str1 contains at least one "cased" character and all of its cased characters are lowercase

str1.isupper()

Returns true if str1 contains at least one "cased" character and all of its cased characters are uppercase

str1.istitle()

Returns true if *str1* is not empty, all of its uppercase characters follow "uncased" characters (spaces, punctuation, etc.), and all of its lowercase characters follow cased characters

Searching

The following methods search for one string inside another. In the descriptions that follow, *startpos* and *endpos* function as they do in slices:

```
str1.startswith(str2[, startpos, [endpos]])
```

Returns true if *str1* starts with *str2*

```
str1.endswith(str2[, startpos, [endos]])
```

Returns true if *str1* ends with *str2*

```
str1.find(str2[, startpos[, endpos]])
```

Returns the lowest index of str1 at which str2 is found, or -1 if it is not found

```
str1.rfind(str2[, startpos[, endpos]])
```

Performs a reverse find: returns the *highest* index where *str2* is found in *str1*, or **-1** if it is not found

```
str1.index(str2[, startpos[, endpos]])
```

Returns the lowest index of str1 at which str2 is found, or ValueError if it is not found

```
str1.rindex(str2[, startpos[, endpos]])
```

Returns the *highest* index of str1 at which str2 is found, or ValueError if it is not found

```
str1.count(str2[, startpos[, endpos]])
```

Returns the number of occurrences of str2 in str1

Replacing

Methods that return a new string with parts of the old string replaced with something else form the basis of a lot of code. The replace method is used particularly frequently, but there are also two other methods that constitute a powerful little facility (though they aren't used all that much):

```
str1.replace(oldstr, newstr[, count])
```

Returns a copy of str1 with all occurrences of the substring oldstr replaced by the string *newstr*; if *count* is specified, only the first *count* occurrences are replaced.

```
str1.translate(dictionary)
```

With dictionary having integers as keys, returns a copy of str1 with any character char for which ord(char) is a key in dictionary replaced by the corresponding value. Exactly what the replacement does depends on the type of the value in the dictionary, as follows:

None

Character is removed from str1

Integer n

Character is replaced by chr(n)

String str2

Character is replaced by str2, which may be of any length

```
str.maketrans(x[, y[, z]])
```

(Called directly through the str type, not an individual string.) Produces a translation table for use with translate more conveniently than manually constructing the table. Arguments are interpreted differently depending on how many there are:

х

x is a dictionary like that expected by translate, except that its keys may be either integers or one-character strings.

x, y

x and y are strings of equal length; the table will translate each character of x to the character in the corresponding position of y.

X, y, z

As with two arguments, plus all characters in the string z will be translated to x (i.e., removed).

Changing case

The methods listed in this section return a new string that is a copy of the original with its characters converted as necessary to a specified case:

str1.lower()

Returns a copy of the string with all of its characters converted to lowercase

Returns a copy of the string with all of its characters converted to uppercase

str1.capitalize()

Returns a copy of the string with only its first character capitalized; has no effect if the first character is not a letter (e.g., if it is a space)

str1.title()

Returns a copy of the string with each word beginning with an uppercase character and the rest lowercase

str1.swapcase()

Returns a copy of the string with lowercase characters made uppercase and vice versa

Reformatting

Each of the methods in this group returns a string that is a copy of the original after applying text formatting operations:

str1.lstrip([chars])

Returns a copy of str1 with leading characters removed. chars is a string that includes the characters to be removed (any combination of them); if it is omitted or None, the default is whitespace.

str1.rstrip([chars])

Returns a copy of *str1* with trailing characters removed. *chars* is a string that includes the characters to be removed (any combination of them); if it is omitted or None, the default is whitespace.

str1.strip([chars])

Returns a copy of str1 with leading and trailing characters removed. chars is a string that includes the characters to be removed (any combination of them); if it is omitted or None, the default is whitespace.

```
str1.ljust(width[, fillchar])
```

Returns str1 left-justified in a new string of length width, "padded" with fillchar (the default fill character is a space).

str1.rjust(width[, fillchar])

Returns str1 right-justified in a new string of length width, "padded" with *fillchar* (the default fill character is a space).

```
str1.center(width[, fillchar])
```

Returns str1 centered in a new string of length width, "padded" with fillchar (the default fill character is a space).

```
str1.expandtabs([tabsize])
```

Returns str1 with all of its tabs replaced by one or more spaces according to *tabsize* (the default is 8) and the column at which the tab occurs on its line.

The format method and function

Many Python programs produce formatted text output. While the methods described in the preceding sections provide a substantial set of operations for manipulating individual strings, arranging the output of several pieces of information together can be a rather intricate project. Python includes a very powerful general formatting facility that can be invoked either through the built-in function called format or through a string method of the same name. These can save you a lot of effort:

```
format(value[, format-specification])
```

Returns a string obtained by formatting value according to the formatspecification; if no format-specification is provided, this is equivalent to str(value)

```
format-specification.format(posargs, ..., kwdargs, ...)
```

Returns a string formatted according to the *format-specification*; any number of positional arguments may be followed by any number of keyword arguments

Format specifications incorporate values where curly braces occur. A pair of braces can contain a number (to indicate a positional argument), a name (to indicate a keyword argument), or neither (to indicate the next positional argument). However, empty braces cannot appear in a format string after braces containing numbers or names. The advantage of using names in format specifications is the same as that of using keyword arguments in function definitions: they describe the role the value plays in the formatted string, and they allow the arguments to be specified in any order.

Given:

```
str1 = 'a string'
```

the following all produce the result '"a string" contains 8 characters':

```
'"{0}" contains {1} characters'.format(str1, len(str1))
'"{}" contains {} characters'.format(str1, len(str1))
'"{string}" contains {length} characters'.format(string=str1,
length=len(str1))
'"{string}" contains {length} characters'.format(length=len(str1),
string=str1)
```

The format method and function can do more than simply replace values. The format specification dictates how each value is formatted within the resulting string. A colon inside a pair of braces begins a format specifier. The complete general structure of a format specifier is as follows:

```
[[fill]align][sign][#][0][width][.precision][type]
```

Some of these you will use only rarely, but when you need them they give you all the control you need over the format of a string. The values for the fill, align, and width fields are listed in Table 3-8. The numeric parts of the specifier are described in Table 3-9, and Table 3-10 gives the options for integer and floating-point type specifications.

Table 3-8. General format specifier fields

Field	Option	Meaning
align	<	Align left (the default)
	>	Align right
	۸	Center
	=	Padding goes between sign and digits (for numeric values only)
fill	not }	Fill character used for a ligning; if the second character of the specifier is not a fill option, the first character is not interpreted as a fill character
width	integer	Minimum field width

Table 3-9. Numeric format specifier fields

Field	Option	Meaning
sign	+	Indicates that signs should be used for both positive and negative numbers
	-	Indicates that a sign should be used only for negative numbers (the default)
	space	Indicates that a leading space should be used for positive numbers
#		Only for binary, octal, or hexadecimal output of integers: prefix the number with Ob, Oo, or Ox, respectively
0	zero-padding	Preceding width, indicates zero-padding; equivalent to alignment = and fill character 0 $$
precision	integer	The number of digits to be displayed after the decimal point of a number formatted with f or F or before and after the decimal point of a number formatted with g or G

Table 3-10. Numeric type specifier fields

Field	Option	Meaning
Types for	b	Binary (base 2)
integers	С	Character: converts the integer to the corresponding Unicode character
	d	Decimal digit (base 10; the default)
	0	Octal (base 8)
	Х	Hexadecimal (base 16), using lowercase letters a—f
	Χ	Hexadecimal (base 16), using uppercase letters A–F
	n	Like d but uses the locally appropriate number separators
Types for	е	Exponent notation (scientific notation using e for the exponent)
floating-	E	Same as e but uses uppercase E for the exponent
point and decimal	f	Fixed point
values	F	Same as f
	g	"General" format (the default): prints the number as fixed point unless it is too large, in which case it switches to e
	G	Same as g but uses E
	n	Like g but uses the locally appropriate number separators

Ranges

A range represents a series of integers. We'll see some uses for ranges later in this chapter and also in the next. They may not look like much, but their use is quite fundamental to Python programming.

With one argument, range(stop) creates a range representing the integers from 0 up to but not including stop. With two arguments, range(start, stop) creates a range representing the integers from *start* up to but not including *stop*. (If *stop* is less than or equal to start, the range will be empty.) With three arguments, range(start, stop, step) creates a range representing the integers from start up to but not including stop, in increments of step. Normally, if step is negative, stop will be less than start. Note the similarity between ranges and slice expressions. Here are some usage examples:

```
>>> range(5)
range(0, 5)
>>> set(range(5))
\{0, 1, 2, 3, 4\}
>>> range(5, 10)
range(5, 10)
>>> set(range(5, 10))
\{8, 9, 5, 6, 7\}
>>> range(5, 10, 2)
range(5, 10, 2)
>>> set(range(5, 10, 2))
```

```
\{9, 5, 7\}
>>> set(range(15, 10, -2))
{15, 13, 11}
>>> set(range(0, -25, -5))
\{0, -15, -5, -20, -10\}
```

Tuples

A tuple is an immutable sequence that can contain any type of element. A very common use of tuples is as a simple representation of pairs such as x, y coordinates. For example, the built-in function divmod(x,y) returns a two-element tuple containing the quotient and the remainder of the integer division of x by y. Tuples are also the natural form for record-oriented data obtained from an external source, such as a row in a database.

Tuple syntax

Tuples are written as comma-separated series of items surrounded by parentheses. This leads to the only ambiguity in Python's syntax: since parentheses are used to group expressions, a parenthesized expression such as (3 * 5) cannot be read as a one-element tuple. To get around this problem, one-element tuples must be written with a comma after their single element. (Commas may always follow the last element of any sequence display.) An empty pair of parentheses creates an empty tuple. Here are some examples:

```
>>> ('TCAG', 'UCAG')
                                 # a two-element tuple
('TCAG', 'UCAG')
>>> ('TCAG',)
                                 # a one-element tuple
('TCAG',)
>>> ()
                                 # an empty tuple
()
>>> ('TCAG')
                                 # not a tuple!
```

Given a sequence as an argument, the tuple function creates a tuple containing the elements of the sequence. Remember, a string is considered a sequence of one-character strings in contexts like this:

```
>>> tuple('TCAG')
('T', 'C', 'A', 'G')
>>> tuple(range(5,10))
(5, 6, 7, 8, 9)
```

Tuple packing and unpacking

The righthand side of an assignment statement can be a series of comma-separated expressions. The result is a tuple with those values. This is called *tuple packing*:

```
>>> bases = 'TCAG', 'UCAG'
                               # a two-element tuple
>>> bases
('TCAG', 'UCAG')
```

Multiple expressions in a return statement are also packed into a tuple, so this is a very convenient way for a function to return more than one value. Example 3-2 illustrates a multiple return from a function. It uses the recognition site function defined in the preceding chapter to simulate the enzyme's cut.

Example 3-2. Returning multiple values from a function

```
def restriction cut(base seq, recognition seq, offset = 0):
    ""Return a pair of sequences derived from base_seq by splitting it at the first appearance
    of recognition_seq; offset, which may be negative, is the number of bases relative to the
    beginning of the site where the sequence is cut"""
    site = recognition site(base seq, recognition seq)
    return base seq[:site+offset], base seq[site+offset:]
```

A tuple—either explicitly enclosed in parentheses or packed—can be the lefthand side of a simple assignment statement (but not an augmented assignment statement, such as one that uses +=). The righthand side of the assignment statement must be a sequence containing the same number of elements as the tuple on the left, either as an implicitly packed tuple or a single sequence value. For example:

```
>>> DNABases, RNABases = 'TCAG', 'UCAG'
>>> DNABases
'TCAG'
>>> RNABases
'UCAG'
>>>
```

A tuple on the left and a tuple-returning function on the right is a fairly common construction. Sometimes a single name will be assigned to the tuple result, but often a series of names will be assigned to the unpacked values that are returned:

```
>>> aseq1 = 'AAAAATCCCGAGGCGGCTATATAGGGCTCCGGAGGCGTAATATAAAA'
>>> left, right = restriction cut(aseq1, 'TCCGGA')
>>> left
'AAAAATCCCGAGGCGGCTATATAGGGC'
>>> right
'TCCGGAGGCGTAATATAAAA'
```

An interesting consequence of this is that the bindings of two names can be exchanged in one statement. This kind of situation does arise from time to time. This works because all the values on the right are evaluated, then packed into a tuple, and finally unpacked for assignment on the left:

```
>>> left, right = right, left
```

Lists

A list is a mutable sequence of any kind of element. Lists are a highly flexible and widely applicable kind of container—so much so that they could be considered the archetypal collection type. One way of thinking of lists is as mutable tuples, since anything you can do with a tuple you can also do with a list. However, you can do a lot more with lists than you can with tuples.

The syntax for lists is a comma-separated series of values enclosed in square brackets. (Square brackets do not lead to the kind of ambiguity with a one-element list that parentheses do with tuples.)

A few built-in functions return lists. For example, dir(x) returns a list of the names associated with whatever x is. As with help(x), the list will contain more than you probably want to see: it includes all the names beginning with underscores, which are meant to be internal to x's implementation. Later in this chapter (Example 3-4), we'll define a function called dr that filters the result of calling dir to omit the private names.

There are many operations and methods that modify lists. Even slicing can be used to change what a list contains. Lists are often used as a way of producing a modified copy of an immutable sequence. First, a new list is created by calling list with an immutable sequence as its argument; then the list is modified, and finally, the type of the immutable sequence is called with the modified list as its argument. As you'll see in the next chapter, another common use of lists is to accumulate values.

Bytearrays are essentially lists with elements restricted to the integers 0 to 255 inclusive. All the operations and methods that you can use for modifying a list can also be used to modify a bytearray.

Statements that modify lists

An index expression designates a particular element of a sequence. A slice specifies a subsequence. Index and slice expressions of lists and bytearrays—the two mutable sequence types—can appear on the lefthand side of an assignment statement. What this means is that the element or subsequence represented by the index or slice expression is replaced by the value of the expression on the assignment statement's righthand side. The change is made to the list itself, not to a copy (the way it would be with string operations).

Slice assignments can express many kinds of list manipulations, as shown in Table 3-11. Some can even change the size of the list. It is well worth studying their varieties.

Table 3-11. List modification assignments

Assignment expression	Result
<pre>lst[n] = x</pre>	Replaces the n^{th} element of $1st$ with x
lst[i:j] = coll	Replaces the $i^{ ext{th}}$ through $j^{ ext{th}}$ elements of lst with the elements of $coll$
<pre>lst[i:j] = any_empty_collection</pre>	Deletes the i^{th} through j^{th} elements of lst (an important special case of $lst[i:j] = seq$)
<pre>lst[i:j:k] = coll</pre>	Replaces the elements of $1st$ designated by the slice with the elements of $coll$, whose length must equal the number of elements designated by the slice
<pre>lst[n:n] = coll</pre>	Inserts the elements of $col1$ before the n^{th} element of $1st$
lst[len(lst):len(lst)] = [x]	Adds x to the end of 1st

Assignment expression	Result
<pre>lst[len(lst):len(lst)] = coll</pre>	Adds the elements of $coll$ at the end of lst
1st += coll	
lst[:] = coll	Replaces the entire contents of $1st$ with the elements of $coll$

There is a simple statement that can remove elements: del, for "delete."

STATEMENT Deletion The del statement removes one or more elements from a list or bytearray. del lst[n]# remove the nth element from lst del lst[i:j] # remove the ith through ith elements from lst $del\ lst[i:j:k]\ \#\ remove\ every\ k\ elements\ from\ i\ up\ to\ j\ from\ lst$

List modification methods

Table 3-12 shows the methods that change a list. These methods are unusual in that they actually change the list itself, rather than producing a modified copy as would similar methods of other types. They are also unusual because—with the exception of pop—they do not return a value (i.e., they return None).

Table 3-12. List modification methods

Method	Result
<pre>1st.append(x)</pre>	Adds x to the end of 1st
<pre>lst.extend(x)</pre>	Adds the elements of x at the end of $1st$
<pre>lst.insert(i, x)</pre>	Inserts x before the $i^{ ext{th}}$ element of $1st$
<pre>lst.remove(x)</pre>	Removes the first occurrence of x from $1st$; an error is raised if x is not in $1st$
lst.pop([i])	Removes the i^{th} element from $1st$ and returns it; if i is not specified, removes the last element
<pre>lst.reverse()</pre>	Reverses the list
<pre>lst.sort([reverseflag] [, keyfn])</pre>	Sorts the list by comparing elements or, if $keyfn$ is included in the call, comparing the results of calling $keyfn$ for each element; if $reverseflag$ is true, the ordering is reversed; $keyfn$ and $reverseflg$ must be specified as keyword arguments, not positionally a

Atypically, the optional arguments to sox t may not be supplied positionally. Either or both may be supplied, but only as keyword arguments. The parameter reverse is simply a flag that controls whether the list is sorted in increasing or decreasing order. The keyfn parameter is explained at the end of the chapter, in the section "Functional Parameters" on page 89.

Sets, frozensets, strings, tuples, and lists can be concatenated with other values of the same type, and the result is a new value of the same type. List modifications are different. No new list is created; instead, the contents of the original list are changed. This is particularly evident in the different results obtained by concatenating two lists as

opposed to using the extend method, as demonstrated in the following interaction and Figure 3-1:

```
>>> list1 = [1,2,3]
>>> list2 = [4,5]
>>> list1 + list2
                               # concatenation
                               # produces a new list
[1, 2, 3, 4, 5]
>>> list1
[1, 2, 3]
                               # while list1 remains unchanged
>>> list2
                               # as does list2
[4, 5]
>>> list1.extend(list2)
                               # extension
>>> list1
[1, 2, 3, 4, 5]
                               # modifies list1
>>> list2
                               # but not list2
[4, 5]
```

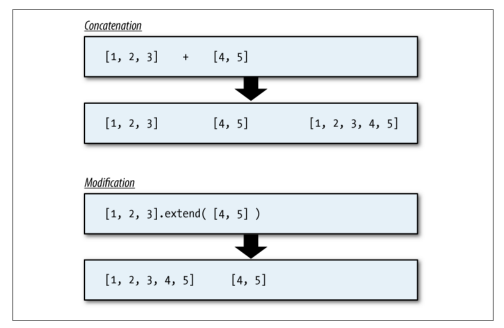


Figure 3-1. List concatenation versus list modification



Augmented assignment of one list to another modifies the list on the lefthand side of the statement.

Sequence-oriented string methods

There are some important string methods that we haven't looked at yet; they were not included earlier in the chapter because we hadn't yet discussed the types of sequences they take as arguments or return as results. In practice the sequence arguments are

usually lists, but in principle they can be any kind of sequence. Remember too that when a sequence argument is expected, a string is interpreted as a sequence of onecharacter strings. Here are the remaining string methods:

string.splitlines([keepflq])

Returns a list of the "lines" in string, splitting at end-of-line characters. If keepflq is omitted or is false, the end-of-line characters are not included in the lines; otherwise, they are.

string.split([sepr[, maxwords]])

Returns a list of the "words" in string, using sepr as a word delineator. In the special case where sepr is omitted or is None, words are delineated by any consecutive whitespace characters; if maxwords is specified the result will have at most maxwords +1 elements.

string.rsplit([sepr[, maxwords]])

Performs a reverse split: same as split except that if *maxwords* is specified and its value is less than the number of words in *string* the result returned is a list containing the *last maxwords*+1 words.

sepr.join(seg)

Returns a string formed by concatenating the strings in seq separated by sepr, which can be any string (including the empty string).

string.partition(sepr)

Returns a tuple with three elements: the portion of *string* up to the first occurrence of sepr, sepr, and the portion of string after the first occurrence of sepr. If sepr is not found in *string*, the tuple is (*string*, '', '').

string.rpartition(sepr)

Returns a tuple with three elements: the portion of *string* up to the last occurrence of sepr, sepr, and the portion of string after the last occurrence of sepr. If sepr is not found in *string*, the tuple is ('', '', *string*).

Mappings

A mapping is a mutable unordered collection of key/value pairs.† Computer scientists use a number of other names to refer to data structures implementing mappings, including associative arrays, lookup tables, and hash tables. A physical dictionary is a realworld example of a mapping: given a word, you get a definition. Figure 3-2 illustrates the concept.

[†] The term "mapping" comes from mathematics, where it represents a function from a "domain" of values to a "range" of values.

key4 key1	value4
key1	value1
key3	value3
key2	value2

Figure 3-2. A dictionary

Dictionaries

Dictionaries have many applications in programming. There are reasons for Python to have sets, frozensets, bytes, bytearrays, and ranges, but you could easily write most of your programs using just strings, lists, and dictionaries. There is just one mapping type in Python: dict, for "dictionary." Like other collection types, dict can be called with a collection argument to create a dictionary with the elements of the argument. However, those elements must be tuples or lists of two elements—a key and a value:

Because they are so frequently used, Python provides a notation for dictionaries similar to that of sets: ‡ a comma-separated list of key/value pairs enclosed in curly braces, with each key and value separated by a colon. You can put spaces before, after, or both before and after the colon. They are printed with no space before and a space after the colon, so you might want to choose that style. Empty braces create an empty dictionary (not an empty set). The order within the braces doesn't matter, since the dictionary implementation imposes its own order. A simple example demonstrates that:

```
>>> {'A': 'adenine', 'C': 'cytosine', 'G': 'guanine', 'T': 'thymine'}
{'A': 'adenine', 'C': 'cytosine', 'T': 'thymine', 'G': 'guanine'}
```

The keys of a mapping must be unique within the collection, because the dictionary has no way to distinguish different values indexed by the same key. As with sets, the implementation of dict does not allow keys to be instances of mutable built-in types.

[‡] You could consider a set to be a dictionary where the value associated with each key is the key itself; in fact, they are often implemented that way in various languages and libraries.

Even a tuple used as a key may not contain mutable built-in types. Any other kind of object may be used as a key, including numbers, strings, and frozensets.



What happens if the argument added to a set or a key used with a dictionary is a collection with mutable elements? Here's an example:

```
# attempt to create a set
                                      # from a list of two lists
Traceback (most recent call last):
 File "<stdin>", line 1, in <module>
TypeError: unhashable type: 'list'
```

"Hashable" refers to the computation performed as part of storing items in lists and dictionaries so that they can be retrieved extremely efficiently. That's all you need to know about it.

Error Message Vocabulary

Error messages developed for a programming language are phrased using a reasonable technical vocabulary, but not one beginning users—even if they are experienced programmers—will necessarily understand. (Even if you're familiar with the concept that a term refers to, you may not recognize it because you may know it by a different name.) Some error messages are pretty clear; for example, NameError: name 'Non' is not defined. Others are more "opaque"—you'll learn what they mean, but you may not understand why they say what they do.

As you gain experience, you'll develop your own mental dictionary that uses these error messages as keys and has explanations as values. In this case, you just have to learn that "unhashable" refers to the technical limitation that prevents mutable built-in collections from being used as elements of sets or keys of dictionaries. Summaries of some of these translations will be provided at the end of the chapters in which they appear.

Dictionary example: RNA codon translation table

Dictionaries are the natural Python representation for tabular data. For example, they can be used to hold rows obtained from a database table: the table's primary key is the dictionary's key, and the value can be a tuple, with each element of the tuple representing one column of the table. § (There is nothing strange about a dictionary key being part of the value associated with it—in fact, this is quite common.)

Table 3-13 shows the usual RNA codon-to-amino acid translation table. Example 3-3 shows how to represent this as a Python dictionary.

§ If you are not familiar with these database terms, you might want to read the section "Database Tables" on page 360.

Table 3-13. The RNA amino acid translation table

		Second Base			
First Base	U	C	A	G	Third Base
U	Phe	Ser	Tyr	Cys	U
	Phe	Ser	Tyr	Cys	C
	Leu	Ser	stop	stop	A
	Leu	Ser	stop	Trp	G
C	Leu	Pro	His	Arg	U
	Leu	Pro	His	Arg	C
	Leu	Pro	Gln	Arg	A
	Leu	Pro	Gln	Arg	G
A	lle	Thr	Asn	Ser	U
	lle	Thr	Asn	Ser	C
	lle	Thr	Lys	Arg	A
	Met/start	Thr	Lys	Arg	G
G	Val	Ala	Asp	Gly	U
	Val	Ala	Asp	Gly	C
	Val	Ala	Glu	Gly	A
	Val	Ala	Glu	Gly	G

Example 3-3. The RNA translation table as a Python dictionary

```
RNA codon table = {
#
                                                                    Second Base
#
                                                              C
                                                                                                                                            G
# U
          'UUU': 'Phe', 'UCU': 'Ser', 'UAU': 'Tyr', 'UGU': 'Cys', 'UUC': 'Phe', 'UCC': 'Ser', 'UAC': 'Tyr', 'UGC': 'Cys', 'UUA': 'Leu', 'UCA': 'Ser', 'UAA': '---', 'UGA': '---', 'UGC': 'Leu', 'UCG': 'Ser', 'UAG': '---', 'UGG': 'Urp',
                                                                                                                                                                               # UxU
                                                                                                                                                                               # UxC
                                                                                                                                                                               # UxA
                                                                                                                                                                               # UxG
# C
          'CUU': 'Leu', 'CCU': 'Pro', 'CAU': 'His', 'CGU': 'Arg', 'CUC': 'Leu', 'CCC': 'Pro', 'CAC': 'His', 'CGC': 'Arg', 'CUA': 'Leu', 'CCA': 'Pro', 'CAA': 'Gln', 'CGA': 'Arg', 'CUG': 'Leu', 'CCG': 'Pro', 'CAG': 'Gln', 'CGG': 'Arg',
                                                                                                                                                                               # CxU
                                                                                                                                                                                # CxC
                                                                                                                                                                               # CxA
                                                                                                                                                                               # CxG
# A
          'AUU': 'Ile', 'ACU': 'Thr', 'AAU': 'Asn', 'AGU': 'Ser', 'AUC': 'Ile', 'ACC': 'Thr', 'AAC': 'Asn', 'AGC': 'Ser', 'AUA': 'Ile', 'ACA': 'Thr', 'AAA': 'Lys', 'AGA': 'Arg', 'AUG': 'Met', 'ACG': 'Thr', 'AAG': 'Lys', 'AGG': 'Arg',
                                                                                                                                                                               # AxU
                                                                                                                                                                               # AxC
                                                                                                                                                                               # AxA
                                                                                                                                                                               # AxG
          'GUU': 'Val', 'GCU': 'Ala', 'GAU': 'Asp', 'GGU': 'Gly', 'GUC': 'Val', 'GCC': 'Ala', 'GAC': 'Asp', 'GGC': 'Gly', 'GUA': 'Val', 'GCA': 'Ala', 'GAA': 'Glu', 'GGA': 'Gly',
                                                                                                                                                                               # GxU
                                                                                                                                                                               # GxC
                                                                                                                                                                               # GxA
```

```
'GUG': 'Val', 'GCG': 'Ala', 'GAG': 'Glu', 'GGG': 'Gly'
                                                           # GxG
}
```

Had we used lists, we would have had to write code to search through the list for a codon whose translation we needed. With dictionaries the task is trivial, as you can see in Example 3-4.

Example 3-4. RNA codon lookup from a dictionary

```
def translate RNA codon(codon):
   return RNA codon table[codon]
```

The nicely formatted, organized dictionary in Example 3-3 was constructed manually. Example 3-5 shows what this dictionary looks like when printed in the interpreter with a 75-column window, as in this book's code examples. This is really just one long line, with "wrapping" at the window edge. Since comments are ignored on input, they aren't part of the output. Notice that the key/value pairs in the dictionary Python constructs from the input appear in an arbitrary order—dictionaries are unordered data structures.

Example 3-5. Dictionary representing the RNA codon table

```
>>> RNA codon table
{'CUU': Leu', 'UAG': '---', 'ACA': 'Thr', 'AAA': 'Lys', 'AUC': 'Ile', 'AAC
': 'Asn', 'AUA': 'Ile', 'AGG': 'Arg', 'CCU': 'Pro', 'ACU': 'Thr', 'AGC': 'S
er', 'AAG': 'Lys', 'AGA': 'Arg', 'CAU': 'His', 'AAU': 'Asn', 'AUU': 'Ile',
'CUG': 'Leu', 'CUA': 'Leu', 'CUC': 'Leu', 'CAC': 'His', 'UGG': 'Trp', 'CAA'
: 'Gln', 'AGU': 'Ser', 'CCA': 'Pro', 'CCG': 'Pro', 'CCC': 'Pro', 'UAU': 'Ty
r', 'GGU': 'Gly', 'UGU': 'Cys', 'CGA': 'Arg', 'CAG': 'Gln', 'UCU': 'Ser', 'GAU': 'Asp', 'CGG': 'Arg', 'UUU': 'Phe', 'UGC': 'Cys', 'GGG': 'Gly', 'UGA':
  '---', 'GGA': 'Gly', 'UAA': '---', 'ACG': 'Thr', 'UAC': 'Tyr', 'UUC': 'Phe
, 'UCG': 'Ser', 'UUA': 'Leu', 'UUG': 'Leu', 'UCC': 'Ser', 'ACC': 'Thr', 'U
CA': 'Ser', 'GCA': 'Ala', 'GUA': 'Val', 'GCC': 'Ala', 'GUC': 'Val', 'GGC':
'Gly', 'GCG': 'Ala', 'GUG': 'Val', 'GAG': 'Glu', 'GUU': 'Val', 'GCU': 'Ala'
, 'GAC': 'Asp', 'CGU': 'Arg', 'GAA': 'Glu', 'AUG': 'Met', 'CGC': 'Arg'}
```

The way complicated combinations of collections are printed in the interpreter generally does not reveal the structure of the data. To obtain a function that will help you see the structure of your data, you should include the following line in your Python files:

```
from pprint import pprint
```

This imports the function pprint from the module pprint—short for "pretty-print" and adds it to your namespace so that you can call it as a function without the module prefix. You can make the function even easier to use if you import it as a shorter alias:

```
from pprint import pprint as pp
```

The output will vary with your window width, which you can change in IDLE by dragging the lower-right corner. On the General tab of IDLE's Preferences window, you can also specify the size for the interpreter window when IDLE starts up.

This imports the function but calls it pp in your namespace. Here is what RNA codon table looks like when pretty-printed (to save space, only the first few and last few lines are shown):

```
>>> pp(RNA codon table)
{'AAA': 'Lys', 'AAC': 'Asn',
 'AAG': 'Lys',
 'AAU': 'Asn',
 'ACA': 'Thr',
 # . . . output deleted here
 'UUG': 'Trp',
'UUT': 'Cys',
 'UUA': 'Leu',
 'UUC': 'Phe',
 'UUG': 'Leu',
 'UUU': 'Phe'}
```

This output goes to the other extreme, printing each dictionary entry on a separate line so that the full output takes up 64 lines. Doing this reveals something interesting that you might not have noticed if several key/value pairs were printed on each line: pprint.pprint orders the dictionary entries alphabetically by their keys. Thus, whereas on input the bases were ordered at each position by the usual TCAG, the order in the output was ACGT. We'll see in the next chapter how to write a function that prints the codon table the way it was organized on input.

Dictionary operations

An interesting ambiguity arises when talking about a value being "in a dictionary." Dictionaries consist of key/value pairs, so are we looking at the keys or the values? Python's answer is "the keys": the membership operators in and not in test whether a key is present; max and min compare keys; and all and any test keys.

The syntax for getting and setting the value associated with a key is simple. Because dictionaries are unordered, there is no equivalent to the flexible slicing operations of sequences. Table 3-14 summarizes the index-based dictionary expressions.

Table 3-14. Dictionary indexing and removal

Expression	Result
d[key]	The value associated with key ; an error is raised if d does not contain key
d[key] = value	Associates $value$ with key , either adding a new key/value pair or, if key was already in the dictionary, replacing its value
d[key] ⋅= value	Augmented assignment, with \cdot being any of +, -, *, /, //, %, and **; an error is raised if d does not have a value for key or if the value is not numeric
del <i>d</i> [<i>key</i>]	Deletes key from the dictionary; an error is raised if d does not contain key

Slice assignment can remove an element from a mutable sequence, so it isn't necessary to use del statements with mutable sequences (though they can make your code clearer). Dictionary assignment, however, can only add or replace an element, not remove one. That makes del statements more important with dictionaries than with sequences. While there are dict methods that remove elements from a dictionary, del statements are more concise.

Dictionary methods

Because items in a mapping involve both a key and an associated value, there are methods that deal with keys, values, and key/value pairs. In addition, there are methods that perform the equivalent of the dictionary operations but with more options. There is also a method for adding the key/value pairs from another dictionary, replacing the values of any keys that were already present. Explanations of the details of these operations are found in Table 3-15.

Table 3-15. Dictionary methods

Expression	Result
<pre>d.get(key[, default_value])</pre>	Like $d[key]$, but does not cause an error if d does not contain key ; instead, it returns $default_value$, which, if not provided, is None
<pre>d.setdefault(key[, default_value])</pre>	Like d[key] if key is in d; otherwise, adds key with a value of default_value to the dictionary and returns default_value (if not specified, default_value is None)
<pre>d.pop(key[, default_value])</pre>	Like $deld[key]$, but does not cause an error if d does not contain key ; instead, it returns $default_value$, which, if not provided, is None
d1.update(d2)	For each key in $d2$, sets $d1[key]$ to $d2[key]$, replacing the existing value if there was one
d.keys()	Returns a special sequence-like object containing the dictionary's keys
<pre>d.values()</pre>	Returnsaspecialsequence-likeobjectcontainingthedictionary'svalues
<pre>d.items()</pre>	Returns a special sequence-like object containing (key , $value$) tuples for the dictionary's keys

Note that the last three methods return "sequence-like objects": they aren't sequences, but they can be used as if they were in many contexts. If you need a dictionary's keys, values, or items in the form of a list, simply call list with what the corresponding method returns.

Streams

A stream is a temporally ordered sequence of indefinite length, usually limited to one type of element. Each stream has two ends: a source that provides the elements and a sink that absorbs the elements. The term "stream" is apt, conjuring as it does the flow of water into or out of a hose. The more common kinds of stream sources are files, network connections, and the output of a kind of function called a generator. Files and network sources are also common kinds of sinks.

Your input to a command-line shell or the Python interpreter becomes a stream of characters (see Figure 3-3). When Python prints to the terminal, that too is a stream of characters. These examples demonstrate the temporal nature of streams: their elements don't necessary exist anywhere other than in the mechanisms that implement the stream itself. For instance, the keystrokes don't "come from" anywhere—they are events that happen in time. (Stream implementations usually involve some kind of buffer that does hold onto the items coming in or going out, but that is largely an issue of efficiency and doesn't affect the conceptual interpretation of the stream as a data type.)

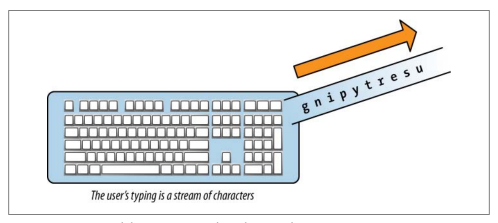


Figure 3-3. A stream of characters issuing from the user's fingers

Files

Usually, the term "file" refers to data stored on media such as a hard disk drive or CD. The term "file" is also used in programming, but with a somewhat different meaning. A Python file is an object that is an *interface* to an external file, not the file itself. File objects provide methods for reading, writing, and managing their instances. The interface is sufficiently natural that the distinction between file objects and the external data they represent can largely be ignored.

The smallest unit of data in external files is, of course, the bit, each representing a 1 or a 0. Bytes are groups of eight bits. They are generally the basic unit of data that moves through the various levels of hardware, operating system software, and programs. Bytes do get grouped into chunks of various kinds throughout the different levels of computer systems, but programmers can usually ignore the details of the larger groupings.

Python's implementation of file objects manages all the details of moving bytes to and from external files. In most respects, a file object is a kind of sequence. Depending on a parameter supplied when an instance is created, the elements of the file object are either bytes or Unicode characters. Some methods treat files as streams of bytes or characters, and other methods treat them as streams of lines of bytes or characters. Figure 3-4 illustrates an input stream and Figure 3-5 an output stream.

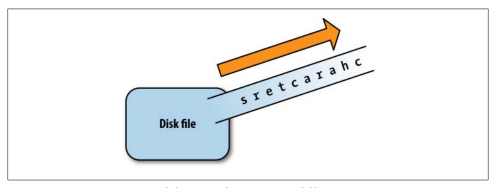


Figure 3-4. Inputting a stream of characters from an external file

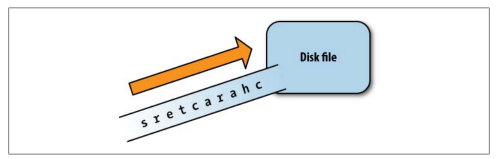


Figure 3-5. Outputting a stream of characters to an external file

Most of the time a file object is a one-way sequence: it can either be read from or written to. It is possible to create a file object that is a two-way stream, though it would be more accurate to say it is a pair of streams—one for reading and one for writing—that just happen to connect to the same external file. Normally when a file object is created, if there was already a file with the same path that file is emptied. File objects can be created to append instead, though, so that data is written to the end of an existing file.

Creating file objects

The built-in function open(path, mode) creates a file object representing the external file at the operating system location specified by the string *path*. The *mode* parameter is a string whose character(s) specify how the file is to be used and its contents interpreted.

The default use is reading, and the default interpretation is text. You can omit the second parameter entirely, in which case the value of *mode* would be 'rt'.

Specifying 'r', 'w', or 'a' without 't' or 'b' is allowable, but the value of mode cannot be just 't' or 'b'. If either of those is specified, 'r', 'w', or 'a' must be explicitly included. If the string contains a plus sign (+) in addition to 'r', 'w', or 'a' the file object will be both readable and writable. The order of the characters within the mode string is immaterial. Content modes are summarized in Table 3-16 and use the modes listed in Table 3-17.

Table 3-16. Content mode values for opening a file

Value	Mode	Interpretation
t	Text (default)	Characters or strings
b	Binary	Bytes

Table 3-17. Use mode values for opening a file

Value	Initial file position	Read?	Write?
r	Beginning (default)	Yes	No
W	Beginning	No	Yes
a	End	No	Yes
r+	Beginning	Yes	No
w+	Beginning	Yes	Yes
a+	End	Yes	Yes

You could use an assignment statement to name the file object created by open, but there's a major problem with this simplistic approach. Due to the way files are implemented both in programming languages and in operating systems, there are various "cleanup" actions that must be performed when a file object is no longer needed. If these actions are not performed after writing with file methods, some characters might not get written to the external file. Other problems can also arise, but that's the most important one.

You could call the method close() to close a file object when it's no longer needed, but if an error occurs while you're using the file, close won't get called, so this won't always work. For this reason Python offers the with statement, which combines naming the file object created by open with a mechanism to ensure that the appropriate cleanup actions are performed. Even if an error occurs during the execution of the statements in a with, the actions necessary to properly close the open file are performed.

STATEMENT

The with Statement

The with statement is used to open and name a file, then automatically close the file regardless of whether an error occurs during the execution of its statements. Like a def statement, a with statement contains an indented block of statements.

```
with open(path, mode) as name:
    statements-using-name
```

More than one file can be opened with the same with statement, as when reading from one and writing to the other.

```
with open(path1, mode1) as name1, open(path2, mode2) as name2, ...:
    statements-using-names
```



In versions of Python before 2.6, the executable first line of a file that uses a with statement must be:

```
from future import with statement
```

File methods

Methods for reading from files include the following:

```
fileobj.read([count])
```

Reads count bytes, or until the end of the file, whichever comes first; if count is omitted, reads everything until the end of the file. If at the end of the file, returns an empty string. This method treats the file as an input stream of characters.

```
fileobj.readline([count])
```

Reads one line from the file object and returns the entire line, including the endof-line character; if count is present, reads at most count characters. If at the end of the file, returns an empty string. This method treats the file as an input stream of lines.

```
fileobj.readlines()
```

Reads lines of a file object until the end of the file is reached and returns them as a list of strings; this method treats the file as an input stream of lines.

File methods for writing to files include the following:

```
fileobj.write(string)
```

Writes *string* to *fileobj*, treating it as an output stream of characters.

```
fileobj.writelines(sequence)
```

Writes each element of *sequence*, which must all be strings, to *fileobj*, treating it as an output stream of lines. Note, however, that although this method's name is intentionally analogous to readlines, newline characters are not added to the strings in *sequence* when they are written to *fileobj*.

Notice that these functions do not provide a location within the sequence to read the characters or lines from or write them to. That's because the location is implicit: the next item in the stream.

In addition, the print function has an optional (keyword) argument, file, that wasn't documented in Chapter 1. When it's included, print writes its output to the value of the file argument rather than to the terminal. The print function can be thought of as a way to convert an object to a string and then send its characters to an output stream.

Example

To manipulate the contents of a file, it is often convenient to just read the entire contents into a string and then manipulate the string instead. This approach runs into trouble with large files, but we'll see ways to fix that in the next chapter. Meanwhile, we'll look at reading FASTA files.#

The first step will be to read the contents of a FASTA file into a string and split that string at every occurrence of '>'. To remind you how str.split works, here's a tiny example. I'll emphasize again the importance of building up your functions from small pieces that you can experiment with to see the behavior:

```
>>> '>id1>id2>id3'.split('>')
['', 'id1', 'id2', 'id3']
```

The list returned by split in this example has a surprising first element. Imagine how difficult it would be to find the source of bugs or erroneous return values if you were writing a 20-line function to fully process a FASTA file. You're better off starting small, with small functions and small input to test them on. There's nothing wrong—and a lot right—with having functions so small that they each do only one significant thing.

We can capture this bit of code directly in a function that substitutes the contents of a file for the string in the example (Example 3-6). The slice at the end of the function eliminates that surprising and useless first element.

Example 3-6. Reading FASTA sequences from a file, step 1

```
def read_FASTA_strings(filename):
    with open(filename) as file:
        return file.read().split('>')[1:]
```

There are some problems with this function, which we'll fix at the end of this chapter: the description line preceding each sequence is part of the sequence string, and the string contains internal newline characters. We will also see many more examples of ways to read data from FASTA files in later chapters.

#FASTA-formatted files are widely used in bioinformatics. They consist of one or more base or amino acid sequences broken up into lines of reasonable size (typically 70 characters), each preceded by a line beginning with a ">" character. That line is referred to as the sequence's *description*, and it generally contains various identifiers and comments that pertain to the sequence that follows.

Since the call to the read method returns the entire contents of the file, we're done with the file after that call. For many situations, though, it's not appropriate to read the entire file. Considerations include:

- The file might be very large, so reading its entire contents could take a long time and use a lot of computer memory.
- The file might contain different sorts of information, each of which needs to be handled differently.
- In many file formats, delimiters separating items aren't as straightforward as they are in FASTA files.

These situations call for using methods that read a restricted amount from the file. The restriction may be in terms of either what it reads—for example, readline—or for how long it reads, by providing an argument to specify the number of characters.

Generators

A generator is an object that returns values from a series it computes. An example is random.randint, as illustrated in Figure 3-6. What's interesting and important about generator objects is that they produce values only on request. Some of the important advantages of generator objects are as follows:

- A generator can produce an infinitely large series of values, as in the case of random.randint; its callers can use as many as they want.
- A generator can encapsulate significant computation with the caller requesting values until it finds one that meets some condition; the computation that would have been needed to produce the rest of the possible values is avoided.
- A generator can take the place of a list when the list is so long and/or its values are so large that creating the entire list before processing its elements would use enormous amounts of memory.

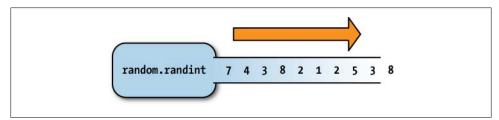


Figure 3-6. A function generating a stream of numbers

The local names of an ordinary function are rebound each time the function is called. The challenge a language faces in implementing generators is that the values of the local names must be retained between calls so that they can be used to compute the next value. (That's how random.randint works, for example.) To fully understand how generators are used, you'll need to learn about some of the statements introduced in the next chapter, but they are mentioned here because of their stream-like behavior.

A value is obtained from a generator by calling the built-in function next with the generator object as its argument. The function that produced the generator object resumes its execution until a yield statement is encountered. At that point, the value of the yield is returned as the value of next. The values of parameters and names assigned in the function are retained between calls. In some simple uses, this is the main purpose of using a generator.

STATEMENT

Yielding a Value from a Generator

A function definition that uses a yield statement in place of return creates a new generator object each time it is called. The generator object encapsulates the function's bindings and code, keeping them together for as long as the object is in use.

yield value

What happens if next is called on a generator when the generator has no more values to produce? The function takes an optional second parameter that specifies the value to return when there are no more values:

next(generator[, default])

Gets the next value from the *generator* object; if the generator has no more values to produce, returns *default*, raising an error if no default value was specified

Collection-Related Expression Features

We've completed our introduction to Python's collection data types. Next, we'll turn to some powerful expression constructs that work with collections.

Comprehensions

A comprehension creates a set, list, or dictionary from the results of evaluating an expression for each element of another collection. Each kind of comprehension is written surrounded by the characters used to surround the corresponding type of collection value: brackets for lists, and braces for sets and dictionaries. We'll start with basic list comprehensions, then look at some variations; finally, we'll explore some examples of set and dictionary comprehensions.



The core of a comprehension is an expression evaluated for each element of a collection. It really must be an expression: statements are not permitted. However, in most situations where you want to use a statement in a comprehension, you can call either a function defined for that purpose or an ad hoc lambda expression, discussed a bit later in this chapter. Function calls are expressions, so they can be used in comprehensions. (Yes, function calls, being expressions, can also be used as statements, but they are still expressions.)

List comprehensions

The simplest form of list comprehension is:

```
[expression for item in collection]
```

This produces a new list created by evaluating the expression with each element of the collection. List comprehensions have wide applicability and can often reduce the complexity of code, making it easier to read and write. You'll probably find them strange at first, but once you're comfortable with them you'll see how powerful and convenient they are. Note that the *expression* must be just that—an expression. Statements are not allowed, but since function calls are expressions they are allowed.

Example 3-7 is yet another rewrite of our validate base sequence function. The expression in this example is base in valid bases and the sequence is base sequence.upper(). (The expression is shown within parentheses to highlight its role, but the parentheses aren't necessary.) The list generated by the comprehension contains only Boolean values. That list is then passed to the built-in function all to check whether or not all the values are true.

Example 3-7. Validating base sequences using a list comprehension

```
def validate base sequence(base sequence, RNAflag = False):
   valid bases = 'UCAG' if RNAflag else 'TCAG'
   return all([(base in valid bases)
               for base in base sequence.upper()])
```

Next, we'll use the functions random base and random codon, which we defined in the previous chapter, to generate a random base sequence. (The function definitions are repeated in Example 3-8 for convenience.) This list comprehension is atypical in that the expression doesn't use the collection items at all—instead, it uses range to determine how many times the expression should be evaluated (i.e., how many codons should go in the list). The expression itself is independent of the numbers generated.

```
Example 3-8. Generating a random list of codons
```

```
from random import randint
def random base(RNAflag = False):
    return ('UCAG' if RNAflag else 'TCAG')[randint(0,3)]
```

```
def random codon(RNAflag = False):
   return random base(RNAflag) + random base(RNAflag) + random base(RNAflag)
def random codons(minlength = 3, maxlength = 10, RNAflag = False):
    """Generate a random list of codons (RNA if RNAflag, else DNA)
   between minlength and maxlength, inclusive""
   return [random codon(RNAflag)
            for n in range(randint(minlength, maxlength))]
```

Often, code makes more sense when it's read from the inside out. Let's take apart the pieces of the last definition to see how it works. We'll assign names to the default values of the parameters so we can evaluate the expressions interactively rather than calling the function:

```
>>> minlength = 3
>>> maxlength = 10
>>> RNAflag = True
```

Next, we name the result of the call randint:

```
>>> randnum = randint(minlength, maxlength)
>>> randnum
4
```

Then we show how that can be used as the collection of a list comprehension:

```
>>> [n for n in range(randnum)]
[0, 1, 2, 3]
```

Finally, we make the expression generate a random codon for each of those integers (ignoring the values of the integers):

```
>>> [random codon(RNAflag) for n in range(randnum)]
['ACT', 'AGG', 'GAG', 'TCT']
```

Of course, since this is all based on a random number, you won't necessarily get the same results if you try these examples yourself.

We can take this example a step further, as shown in Example 3-9. We already have a translate RNA codon function, which we developed in the section "Dictionaries" on page 67. We can use that to define a function that translates a collection of codons. We could also add the code to the random codons function, but as explained earlier, it is often best to leave small function definitions as they are. You can then use them in the construction of other functions, rather than trying to pack all the desired capabilities into one function.

Example 3-9. Translating random base sequences

```
def random codons translation(minlength = 3, maxlength = 10):
    """Generate a random list of codons between minlength and maxlength, inclusive"""
   return [translate RNA codon(codon) for codon in
            random codons(minlength, maxlength, True)]
```

Now let's run a few calls to see what the results look like:

```
>>> random codons translation()
['Ser', 'Gly', 'Ile']
>>> random_codons_translation()
['Ser', 'Leu', 'Ala', 'Leu', 'Asn', 'Val', 'Lys', 'Pro', 'His', 'Tyr']
```

Next, let's revisit our primitive FASTA reader. The beginning of each string in the list returned by the definition in Example 3-6 contains information about a base sequence. The rest of the string represents the base sequence itself. Suppose we want to split the description from a base sequence returned from the function read FASTA strings, as defined in Example 3-6. There happens to be a string method that does just what we need. Given a string string and another string sepr, the call string.partition(sepr) returns a tuple with three elements: the part of string up to the first appearance of sepr, sepr, and the part of string after sepr. Calling partition with an argument of '\n' will split the description from the base sequence.

We can call partition on each string produced by a call to read FASTA strings and use a list comprehension to collect the results. We could add the necessary code to read FASTA strings, but we'll adhere to our recommendation of having each function do only one significant thing. Besides, read FASTA strings is a function that is perfectly good in its own right and that might have uses beyond the current example. Instead, we'll define a new function, read FASTA entries (Example 3-10).

```
Example 3-10. Reading FASTA sequences from a file, step 2
```

```
def read FASTA entries(filename):
   return [seq.partition('\n') for seq in read FASTA strings(filename)]
```

This function returns a list of the sequences from a FASTA file, each one a tuple of the form (information, '\n', sequence). (Remember to use pprint.pprint to help you understand the structure of results that are this complicated.) Next, we need to remove the newline characters from within each sequence string. Again, we'll define a new function: it will use str.replace and another list comprehension. We'll take this opportunity to discard the useless '>' that begins each description. Example 3-11 shows the new function, read FASTA sequences.

Example 3-11. Reading FASTA sequences from a file, step 3

```
def read FASTA sequences(filename):
                                                                # ignore the initial '>'
    return [[seq[0][1:],
              seq[2].replace('\n', '')]
                                                                # delete newlines
             for seq in read FASTA entries(filename)]
```

When the value used in the comprehension—seq in this case—is a sequence, it can be unpacked into multiple names, which helps make the code clearer. Compare Example 3-12 with Example 3-11. It's not a big difference, but it can be helpful in more complicated examples. Naming the elements of the sequence saves you from having to figure out what the elements were when you come back to the code later.

```
def read FASTA sequences(filename):
   return [(info[1:], seq.replace('\n', ''))
            for info, ignore, seq in
                                                              # ignore is ignored (!)
            read FASTA entries(filename)]
```

This is probably all beginning to look a bit rich. Play with the code by trying out its bits separately, as demonstrated earlier. You'll have reached a key stage in your Python education when you can look at a function like read FASTA sequences in Example 3-12 and see more or less immediately what it is doing. In the meantime, it can be very helpful to work through functions like this by entering them into the interpreter line by line and examining the results.

The description lines of FASTA files generally contain vertical bars to separate field values. Example 3-13 defines a new function that calls read FASTA sequences, then uses str.split to return a list of field values for the description instead of just a string.

```
Example 3-13. Reading FASTA sequences from a file, step 4
```

```
def read FASTA sequences and info(filename):
   return [[seq[0].split('|'), seq[1]] for seq in
            read FASTA sequences(filename)]
```

Notice how we continue to keep each function small. When building a "vocabulary" of such functions, you don't know which ones you or other programmers will ultimately need. In any case, it's easier to leave the working definitions alone and build on top of them than it is to keep changing already working functions.

Altogether, the functions we developed to read sequences from a FASTA file do the following:

- Split the file contents at '>' to get a list of strings representing entries
- Partition the strings to separate the first line from the rest
- Remove the useless '>' from the resulting triples
- Remove the newlines from the sequence data
- Split the description line into pieces where vertical bars appear

Each sequence in the result returned is represented by a two-element list. The first element is a list of the segments of the description, and the second is the whole sequence with no newline characters.

What would it look like to do this all at once? A combined definition is shown in Example 3-14.

Example 3-14. Reading FASTA sequences with one compact function

```
def read FASTA(filename):
   with open(filename) as file:
        return [(part[0].split('|'),
```

```
part[2].replace('\n', ''))
for part in
[entry.partition('\n')
for entry in file.read().split('>')[1:]]]
```

This isn't terrible, but it's near the limit of what you could hope to readily comprehend even once you're comfortable with code like this. You'd probably try to understand it by looking at each piece individually to see what it does, so why not just define each piece as a separate function that names what it does? Doing so makes it easier to read, test, and fix your code.

If ever you do want to produce an all-in-one function like the one in Example 3-14, you should still develop it piece by piece, working from the inside out. Follow the same steps we followed in developing the sequence of separate functions, making one change to the function at a time. It's true that using four separate functions requires a few more lines of code, but they are much more readable than the ensemble of maneuvers in the all-in-one definition. Also, some of them might prove useful in situations that call for less processing of each entry.

Yet another approach is shown in Example 3-15, which defines the function as a series of steps, naming the results of each. This does help to clarify what each piece is doing and can help with debugging, but it's a style more appropriate for programming in older languages. Python's expressiveness and clarity make assignment statements superfluous in many situations. That said, you could write the function using a lot of assignment statements as you are developing. Then, once you've got the definition working, you could remove them and pack everything together into more complex expressions.

Example 3-15. Reading FASTA entries from a file, binding names

```
def read FASTA(filename):
    with open(filename) as file:
         contents = file.read()
                                                                          # only statement inside the with
    entries = contents.split('>')[1:]
                                                                          # skip blank first entry
    partitioned entries = [entry.partition('\n') for entry in entries]
    pairs = [(entry[0], entry[2]) for entry in partitioned entries] # omit '>'
    pairs2 = [(pair[0], pair[1].replace('\n', '')) for pair in pairs]
result = [(pair[0].split('|'), pair[1]) for pair in pairs2]
    return result
```

Finally, there's nothing to prevent you using an intermediary style. For instance, in Example 3-15 there's no reason to have one step that removes the newlines from the second element of the pairs and a second step that splits the first element at vertical bars. These actions could be merged into one step:

```
[(pair[0].split('|'), pair[1].replace('\n', '')) for pair in pairs]
```

Set and dictionary comprehensions

Set comprehensions work like list comprehensions. They are enclosed in braces and produce a set:

```
{expression for item in collection}
```

One important difference from list comprehensions is that the result of a set comprehension doesn't necessarily have as many elements as the collection used by the comprehension. Because it's a set, duplicates accumulated during the comprehension are ignored. Also, the value of expression cannot be an instance of a mutable built-in collection, because that could not be an element of a set.

Dictionary comprehensions are also enclosed in braces, but they are a bit more complicated because they require key/value pairs:

```
{key-expression: value-expression for key, value in collection}
```

Just as when you call dict with a collection argument, the collection in a dictionary comprehension must contain only two-element lists or tuples. As always with dictionaries, the key values may not be mutable collections.

Taking a list of pairs of the form (info-fields, sequence) as produced by read FASTA in Example 3-16 and assuming the first info field is 'gi' and the second the identifier, it's easy to construct a dictionary of sequences indexed by identifier.

Example 3-16. Constructing a dictionary with a comprehension

```
def make indexed sequence dictionary(filename):
   return {info[0]: seq for info, seq in read_FASTA(filename)}
```

Generator expressions

A bit earlier in this chapter we discussed using the vield statement to create a function that returns a new generator object each time it is called. Comprehensions are often a more convenient way to express computations that otherwise would be defined as separate functions, and Python has something similar for generators. A generator expression is syntactically like a list or set comprehension, except that it is surrounded with parentheses and its value is a generator:

```
(expression for item in collection)
```

Since yield is implicit in generator expressions, they are often quite a bit simpler than equivalent generator-producing functions.

Generators look just like equivalent list or set comprehensions. Accessing individual elements of these three collection types requires following different rules, though:

- There is no way to access an individual element of a set.
- There is only one element that can be accessed from a generator, and that is obtained by calling next.
- Any element of a list can be directly accessed with an index expression.

Some collection types can accept any other kind of collection as their argument when called as a function. Thus, set(q) will return a set constructed from all the elements of the generator q, and list(q) will return a list of all the elements of the generator q. Note the similarity to files, another kind of stream: next and list for generators correspond to readline and readlines for files. You can get the next item of the stream, or get them all. However, you cannot ask for a specific item using an index expression.

Generator expressions are useful when the expressions they encompass require substantial computational resources or the collections are large. By using next to get one element of the generated stream at a time and stopping when certain criteria have been fulfilled, we can avoid unnecessary computation. Also, like generator functions, generator expressions can generate infinite streams. (But don't try to call set, list, or tuple on one of those!) Example 3-17 shows how a generator expression could be used to produce amino acid abbreviations from an RNA sequence.

Example 3-17. Generating amino acid translations of codons

```
def aa generator(rnaseq):
    ""Return a generator object that produces an amino acid by translating
    the next three characters of rnaseq each time next is called on it"""
    return (translate RNA codon(rnaseq[n:n+3])
             for n in range(0, len(rnaseq), 3))
```

Since we're ignoring the problem of running out of codons, the definition does not handle the extra base or two that might appear at the end of the string following the last codon. Here's how this works:

```
>>> aagen = aa generator('CCACCGCACCAACAGCGC')
>>> next(aagen, None)
'Pro'
>>> next(aagen, None)
'Pro'
>>> next(aagen, None)
'His'
>>> next(aagen, None)
>>> next(aagen, None)
>>> next(aagen, None)
'Arg'
```

It is an error to call next on a generator that has no more elements to produce, unless a second argument is included in the call. In that case, the second argument is a value to return when the generator is exhausted. It is not an error to call next on an exhausted generator as long as the second argument is included. Consider the following two calls after the preceding series of inputs:

```
>>> next(aagen, None)
                                           # OK, default value None returned
>>> next(aagen)
                                           # Error, since no default value is provided
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
StopIteration
```

StopIteration doesn't look much like the name of an error message, and we haven't even discussed "iteration" yet (we will in the next chapter). Actually, StopIteration is used in the implementation of certain parts of Python, and we'll even use it in some later examples. It's not so much an error as an indication that the end of a stream has been reached. What makes it an error in this case is that the call to next didn't say what to do when the stream was exhausted.

Conditional comprehensions

The value of a comprehension does not necessarily have to include an element corresponding to every element in its collection. One or more tests can be added to determine the elements for which the comprehension expression will be evaluated. This is called *filtering* and is implemented with a *conditional comprehension*:

```
[expression for element in collection if test]
```

Multiple tests can be included as a series of "if" components of the comprehension. Example 3-18 demonstrates the use of a conditional comprehension. The function filters the names returned by dir to eliminate names beginning with underscores.

Example 3-18. Filtering out the underscore names from dir

```
def dr(name):
    """Return the result of dir(name), omitting any names beginning with an underscore"""
    return [nm for nm in dir(name) if nm[0] != ' ']
```

Files often contain different kinds of information on different lines, such as the descriptions and sequences of a FASTA file. This kind of situation is another good candidate for a conditional comprehension. Example 3-19 shows how easy it is to extract just sequence descriptions from a FASTA file and split them into fields at their vertical bars.

Example 3-19. Reading FASTA descriptions from a file

```
def get FASTA descriptions(filename):
   with open(filename) as file:
        return [line[1:].split('|') for line in file if line[0] == '>']
```

Conditional comprehensions work the same way for sets as they do for lists. We could use one to find all the codes in the third position of the description lines of a FASTA file. For example, a file obtained from GenBank will begin with gi, an identifier, gb, and a GenBank accession number. However, the third field of a FASTA file obtained from another source or generated by an application might have a different meaning. Example 3-20 defines a function that looks at the descriptions in a FASTA file and returns a list of all the different codes it finds in the third field.

Example 3-20. Reading FASTA descriptions using set comprehension

```
def get FASTA codes(filename):
   with open(filename) as file:
```

```
if len(line.split('|')) < 3:</pre>
    return []
return {line.split('|')[2] for line in file if line[0] == '>'}
```

Conditional dictionary comprehensions are similar. They use braces and must supply both a key and a value, separated by a colon. In Example 3-20 no provision was made for entries whose descriptions begin with something other than 'gi'. The definition presented in Example 3-21 omits such sequences (which may or may not be a useful thing to do).

Example 3-21. Constructing a selective dictionary

```
def make gi indexed sequence dictionary(filename):
   return {info[1]: seq for info, seq in read FASTA(filename)
           if len(info) >= 2 and info[0] == 'gi'}
```

Even generator expressions can include conditionals. They produce values only for those elements of the collection for which the conditional(s) are true. Example 3-22 shows an interesting use of a conditional generator expression to find the first element in one list that is also in a second.

Example 3-22. Using a generator to find the first common element

```
def first common(list1, list2):
    """Return the first element in list1 that is in list2"""
    return next((item for item in list1 if item in list2), None)
```

Nested comprehensions

Comprehensions can have more than one for in them. When they do, the innermost ranges over its collection for each of the next one out's collection, and so on. It is rare to see more than two for sections in a comprehension, but occasionally three can be useful.

Example 3-23 shows how clear a terse comprehension can be while defining a significant computation. The function returns a list containing every three-character combination of the (unique) characters in its argument. It first converts its argument to a set in order to eliminate any duplicates. The intended use is to generate a list of the strings representing all the codons for a given collection of bases. The default is 'TCAG', but it could be called with 'UCAG'.

Example 3-23. A nested comprehension for generating codons

```
def generate triples(chars='TCAG'):
     ""Return a list of all three-character combinations of unique characters in chars"""
    chars = set(chars)
    return [b1 + b2 + b3 for b1 in chars for b2 in chars for b3 in chars]
```

In fact, the function is entirely general. It will produce a list containing the result of evaluating the + operator with every combination of three values in its argument:

• A string of any length:

```
>>> generate triples('01')
['111', '110<sup>T</sup>, '101', '100', '011', '010', '001', '000']
```

• Any nonstring collection containing one-character strings:

```
>>> generate_triples(('.', '-'))
['---', '---', '---', '---', '.--', '.--', '..-']
```

• Any nonstring collection containing strings of any length:

```
>>> generate_triples({'0', '555'})
['55555555', '5555550', '5550555', '55500', '0555555', '05550', '00555', '000']
```

• Any nonstring collection containing anything that supports +:

```
>>> generate triples(range(2,4))
[6, 7, 7, 8, 7, 8, 8, 9]
```

Functional Parameters

Earlier in this chapter we saw two functions and one method with optional key parameters. How this works requires some discussion.

The parameter "key"

We first encountered the two functions, max and min, in Chapter 1, where functions were introduced. There they were described as taking any number of positional arguments, but these two functions can also be called with a collection as the only positional argument. (That wasn't mentioned before because collections hadn't yet been introduced.) Both forms accept an optional key parameter, which is expected to be a function. The value of the key argument is called on each element of the collection. The values returned by those calls are used to select the maximum or minimum element:

```
>>> max(range(3, 7), key=abs)
>>> max(range(-7, 3), key=abs)
```

Note that although abs was used here to compare the elements of the range, what's returned is the *element* for which abs returned the largest value—not the *value* of calling the key function on the element. For a more interesting illustration, consider a list seq 1ist containing RNA base sequences. We could select the sequence with the lowest GC content by calling min with key = gc content (a function defined in the previous chapter).

Earlier in this chapter, the method list.sort was described. It too can take an optional key parameter. The key parameter works the way it does for min and max—the value of key is called on each element, and the elements of the list are sorted according to the values returned. The sort method uses < to order the elements of the list. If a call does not include a key argument, the elements of the collection being sorted must have

mutually comparable types: for example, a list of numbers or a list of strings, but not a mixed list of numbers and strings. If the call does include a key argument, its value is called for each element and the results are compared using <. Let's look at an example:

```
>>> lst = ['T', 'G', 'A', 'G', 't', 'g', 'a', 'g']
>>> 1st
['T', 'G', 'A', 'G', 't', 'g', 'a', 'g']
>>> lst.sort()
>>> 1st
['A', 'G', 'G', 'T', 'a', 'g', 'g', 't']
```



Sorting 1st in this example did change the order of its elements, but in a way that may surprise you: all uppercase letters are considered "less than" all lowercase letters. This behavior has a long history in computer technology, the details of which are not important; keep it in mind, though, when sorting lists of strings.

The way to have a list sorted according to an order you define is to provide a key function. In this example we can eliminate the distinction between upper- and lowercase letters by providing a key that returns the uppercase (or, equivalently, the lowercase) version of its string argument.



Python 2: The key argument is a function of two arguments, defaulting to cmp, that is called to compare two elements of the collection.

```
>>> lst.sort(key=str.lower)
['A', 'a', 'C', 'c', 'G', 'g', 'T', 't']
```



Another thing you might have noticed is that after sorting using str.lower, each uppercase letter appears before the corresponding lowercase one. This is not due to anything about letter case; for technical reasons it is helpful for sorting functions to be stable, meaning that if two items are equal, they will be in the same order after the sort as they were before it.

One use of sorting by key is to order a list of base sequence strings where some use lowercase and some uppercase (perhaps because they were obtained from different sources, sites, databases, or programs). The default sort does not do what we want:

```
>>> seqs = ['TACCTATACCGGCTA', 'cacctctaccgta', 'AACCTGTCCGGCTA']
>>> seqs.sort()
['AACCTGTCCGGCTA', 'TACCTATACCGGCTA', 'cacctctaccgta']
```

The sequence beginning with 'c' should go between the other two but appears at the end. This is only a small example; imagine what would happen when sorting a large number of sequences, some uppercase and others lowercase (or worse, using mixed case). What we want instead is for sort to ignore case:

```
>>> seqs = ['TACCTATACCGGCTA', 'cacctctaccgta', 'AACCTGTCCGGCTA']
>>> seqs.sort(key=str.lower)
>>> seqs
['AACCTGTCCGGCTA', 'cacctctaccgta', 'TACCTATACCGGCTA']
```

Function objects

The value of the key argument in the preceding examples appeared to be just a name. However, as with any use of names in expressions, it is the value the name is bound to that is actually passed to the function. The value of abs is a built-in function, and the value of str.lower is a method in the class str.

The idea of using a function or method as an argument to another function or method may seem strange, but functions and methods are objects, just like everything else in Python. The def statement creates a function object and names it.

You can type the name of a function in the interpreter to see what kind of object it is (what is printed for built-in functions and methods is somewhat different than what you'll see for ones you define or import, and what is printed for functions differs from what is printed for methods, but the values are all objects):

```
>>> max
<built-in function max>
>>> gc content
<function gc content at 0x74a198>
>>> str.lower
<method 'lower' of 'str' objects>
>>> lower
Traceback (most recent call last):
File "<stdin>", line 1, in <module>
NameError: name 'lower' is not defined
```

The reason for the error in the last example is that there is no function called "lower". Only the str class provides a lower method, so either str or an instance of str must appear before lower, with a dot in between.

Like any value, a function can have multiple names. Binding a name to an existing function is no different than binding a name to an existing object of any other type:

```
>>> fn = gc content
>>> fn('ATCCCGGG')
0.75
```

A function call is a kind of expression. In evaluating the expression Python obtains the function object named in the call, binds its parameters to the arguments in the call, and begins executing the function. What "route" (name) it takes to get to the object has no effect on the process.

Anonymous functions

Passing functions as arguments is not as esoteric as it may seem. There are many situations in which a functional parameter avoids a great deal of repetitive code. Imagine if you had to define a separate function for every different way you might want to find the maximum value in a collection or sort a list. Just as other parameters generalize pieces of computation, so do functional parameters.

Functional parameters are powerful tools, but you still have to write the different functions to pass as arguments. Functions passed as arguments are generally small and simple. Defining a lot of these little functions used solely to pass to other functions gets tedious. Also, definitions used in a subsidiary way like these really shouldn't be as prominent within a file as normal functions—they'd distract the reader and obscure the real structure of the file's code.

Fortunately, Python has a mechanism for defining lightweight functions without using def. These functions don't even have names—they are anonymous. Although they are functions, they are defined by an expression, not a statement. This kind of expression is called a *lambda expression*.* Its syntax is as follows:

```
lambda args: expression-using-args
```

A lambda expression has one or more arguments. The result is an anonymous function that can be used anywhere you would use a named function. The key parameters of min, max, and lst.sort expect functions of one argument, so a lambda passed to one of these must have exactly one argument. The expression can be any kind of expression, including one with function calls, but it must be an expression, not a statement.

Lambda expressions can be passed as arguments, just as if they were named functions. They have a few important benefits, though: they are a little more concise than functions defined with def; they are anonymous, so there's no risk of them being used inappropriately or accidentally elsewhere in your code; and they keep the definition right where it's used, which improves readability. Actually, you can name a lambda expression with an assignment statement, just as you'd name any other value. The effect would be the same as if you had used def to define and name the function. The following two ways of defining the function named fn are in all ways equivalent:

```
def fn (x, y):
    return x*x + y*y
fn = lambda x, y: x*x + y*y
```

In fact, when a lambda expression is passed to another function, it does get named. Just like any value passed as an argument in a function call, the corresponding name in the

^{*} The term "lambda" and the corresponding concept come from formal logic and mathematics.

function's parameter list is bound to the lambda expression. Inside the function definition, there is no difference whatsoever between using a parameter name to call a function that has been defined with def or an anonymous function defined by a lambda expression.

Let's look at an example. Python has two built-in functions called all and any. They each take an iterable as an argument. The function all returns True if all the elements of the iterable are true, while any returns True if any of them are. Arguably, it might have been better for those functions to have been defined to take an optional function to call on each element and then to check the result returned by the function instead of testing the elements themselves. If no function were provided, the default would be to use the element itself.

We can define functions to do this. It is never a good idea to redefine built-in functions, so we must invent two other names: every and some, perhaps. Example 3-24 shows a definition for some that uses a conditional generator. The function calls next once. If there is an *item* in *coll* for which pred(*item*) is true, the generator produces at least one True value. If, however, pred(item) is not true of any item, the generator produces an empty stream and next will use the default value False, as specified in its call.

Example 3-24. Definition of a function with a functional argument

```
def some(coll, pred=lambda x: x):
    """Return true if pred(item) is true for some item in coll"""
    return next((True for item in coll if pred(item)), False)
>>> some([0, '', False])
>>> some([0, '', False])
False
>>> some([0, '', 4])
>>> some([0, 1, 4], lambda x: isinstance(x, int) and x % 2)
                                                                       # odd ints?
True
>>> def odd(n):
       return bool(n % 2)
)>>> some([0, 2, 4], odd)
False
>>> some({'a string', ''}, len)
>>> some({tuple(), ''}, len)
False
```

The last two lines of input illustrate that some can be used on any collection, not just a sequence. The definition of odd and the input before and after it show that the name of a function and a lambda expression have the same effect in a function call.



It is rarely necessary to call bool: anywhere you would use the result of an expression on which you can call bool, you can just use the expression by itself. The definitions of some and odd in this example did use bool because the functions are predicates like all and any; thus, they are expected to always return a Boolean value.

Returning to the example of sorting a list of strings in mixed case, suppose we want to order them by size first, and then alphabetically. We could order them by size easily enough by providing len as the key argument, just as we ordered them alphabetically after converting all of the sequences to lowercase. However, there's no straightforward way to order them by multiple criteria without passing a function defined for that purpose as the key argument.

What would such a function look like? It could be anything that returns a result appropriate for use in a < expression. It turns out that the definition of < when its operands are sequences is to compare their first elements, then their second, and so on, until one is "less than" the other. For example:

```
>>> 1 = [(3, 'abc'), (5, 'ghijk'), (5, 'abcde'), (2, 'bd')]
>>> 1.sort()
>>> 1
[(2, 'bd'), (3, 'abc'), (5, 'abcde'), (5, 'ghijk')]
```

To sort by both length and alphabetically ignoring case, we need a function that returns a pair (tuple or list) whose first element is the sequence's length and its second is the sequence converted to either lower- or uppercase. We could define such a function:

```
def sequence sort key(seq):
    return (len(seq), seq.lower())
```

Then 1.sort(key=sequence sortkey) would do the trick. We probably don't need such a function anywhere else, though, and it's better to use a lambda expression than define a superfluous function:

```
>>> l.sort(key=lambda seq: (len(seq), seq.lower())))
```

It's never *necessary* to use a lambda expression. Once you become familiar with them, though, you'll find them a welcome tool. Even if you don't use them, learning about them will help you better understand the true nature of functions. We'll be using lambda expressions for some important purposes later in the book.

Tips, Traps, and Tracebacks

Tips

• When typing the contents of a collection inside its enclosing punctuation parentheses for tuples, brackets for lists, and curly braces for sets and dictionaries—it's a good idea to always end with an extra comma. That avoids errors if you later add more elements and forget to add the comma after what had previously been the last element. Then, when you do add more elements, make sure you include a comma after the new ones too.

• Keep in mind that there only a few functions that are used with string arguments. Most string operations are method calls; for instance, 'ATTCATT'.count('A').

Language features

- Don't call close on files opened using a with statement; the file will automatically be closed at the end of the statement's execution.
- Practice using comprehensions. They may appear strange at first, but they are an elegant and powerful way to express computations that construct one collection from another, and these are quite common.
- Give generator expressions and generator functions a try.
- Practice using lambda expressions for anonymous function definitions that consist of only an expression.
- Where does Python find files whose names are given as arguments to open? If the argument is an absolute path—i.e., one beginning with a slash—that is the path to the file. Otherwise, the path is relevant to the "current" directory. Determining which directory is current can be confusing. Some of the ways Python determines the current directory are:
 - —When the interpreter is started from a command line, the current directory is the directory that was current in the command-line environment.
 - The directory that is current when IDLE starts is platform-dependent: on OS X, it is the user's Documents folder; on Linux, in might be the home directory or the home/Documents directory; and on Windows, it is the Python installation directory (although, as with all Windows programs, an alias to IDLE can be set up and given a different start directory through its property dialog).
 - —When a file is run from IDLE, the current directory is changed to the directory of the file.
- You can find out what the current directory is by typing the following to the interpreter:

```
import os
os.getcwd()
```

Developing and testing code

- Build programs step by step, keeping function definitions small and testing your code each time you add one or two new ones.
- If you test your code after every change, or every few changes, you'll have a much better idea of what caused a new error than if you write a whole bunch of new code

and try to make it work. Ideally, you should test every new function definition immediately, then test how other functions work when you add calls to the new function to them.

- Use assertion statements liberally to:
 - —Validate arguments.
 - —Validate expected partial results in the middle of function definitions.
 - —Validate results obtained from calls to one of your functions from another.

Why not just use calls to print? There are several reasons:

- —It can be quite a strain—and quite boring—to repeatedly examine similarlooking output as you write and test your code.
- —It is very easy to overlook small problems in the output.
- —The details of formatting output sufficiently to be able to examine it are often needed only for that purpose, in which case it is a waste of time—let Python do the checking with assert statements.

Very common tests to put in assertions include:

- —The expected length of a collection.
- —The range of a numerical value returned from a call to one of your functions (e.g., gc content).
- —The contents of lines read from a file.
- When you do look at collection values—especially as you begin working with collections that contain other collections—remember to use pprint.pprint. You might as well just put this line:

from pprint import pprint as pp

at the beginning of every file to make it easier to use.

 Periodically step back from the details of the code you are developing and look at its overall "shape." Keep an eye out for long functions, repeated code, strings that appear in more than one place, and so on; these are candidates for new definitions and assignments.

Traps

- (value) is simply value with parentheses around it, so it is equal to value, not a one-element tuple. A one-element tuple is written with a trailing comma: e.g., (value,).
- An augmented assignment list1 += list2 modifies list1, whereas list1 + list2 creates a new list.
- The result of file.readline includes a newline character; the lines returned by file.readlines do not.

• In a comprehension that just selects elements of a collection, it is very easy to forget to include the for part, since a comprehension like this looks perfectly natural:

```
[x in lst if x > 0]
```

It should be:

```
[x for x in lst if x > 0]
```

- The result returned by max or min is an element of its collection argument, even if a keyfn argument is provided. The keyfn is used to compute a value for the purpose of comparison; when max or min identifies the appropriate element it returns that element, not the result of calling keyfn on that element.
- The keys, values, and items methods of dict look like they should return lists, but they don't; they return special types of objects, which you will see if you call one of them on a dictionary. These objects are iterables, but they are not in themselves equal to any list. Usually the only thing you do with the results of these methods is use them in a for statement or comprehension. If you do need a list value, call list on the result of the method call, but keep in mind that the order within the list is not predictable.
- Your computer's operating system may be suppressing common file extensions in file browser windows. This can lead to all sorts of problems because you are not seeing the actual name of the filename. Turn off this consumer-oriented feature, as follows:

OSX

In the Finder Preferences, click Advanced (the gear icon) at the top and turn on "Show all File Extensions."

Windows

In a Windows Explorer window, select "Folder Options" from the Tools menu, click the View tab, and uncheck "Hide extensions for known file types."

Unix/Linux

Different desktop environments may do this differently, but you should be able to figure out how to make sure file extensions are always shown in the standard file browser.

 To open a file that contains Unicode characters, it is necessary to include encoding='utf-8' as the third argument to open.

Tracebacks

Representative error messages include:

```
NameError: [len(x) in lst]
    Missing for x before in.
```

SyntaxError: [len(x) in lst if x > 3]

Missing for x before in.

IndexError: string index out of range

For a string of length N, an index (i.e., the value between square brackets) must be in the range -N <= index < N-1.

TypeError: unhashable type 'list'

TypeErrors have many causes and variations. "Unhashable" indicates a violation of the prohibition against using mutable built-in types as elements of sets or keys of dictionaries.

object is not iterable: 'builtin function or method'

A function name was used in place of a function call in the in part of a comprehension; i.e., you forgot the parentheses! This is a common mistake when calling dict.keys(), dict.values(), and dict.items().

Control Statements



This chapter's material is rich and intellectually challenging. Don't give up if you start to feel lost (but do review it later to make sure you have absorbed it all). This chapter, together with the next, will complete our introduction to Python. To help you understand its contents, the chapter ends with some extended examples that reiterate the points made in its shorter examples. The rest of the book has a very different flavor.

Chapters 1 and 2 introduced *simple statements*:

- Expressions, including function calls
- Assignments
- Augmented assignments
- Various forms of import
- Assertions
- return
- yield (to implement generators)
- pass

They also introduced the statements def, for defining functions, and with, to use with files.* These are *compound statements* because they require at least one indented statement after the first line. This chapter introduces other compound statements. As with def and with, the first line of every compound statement must end with a colon and be followed by at least one statement indented relative to it. Unlike def and with statements, though, the other compound statements do not name anything. Rather, they determine the order in which other statements are executed. That order is traditionally

^{*} The with statement is more general than how it was described in Chapter 2: it actually does not need to name the object of the with in an as portion of the statement, and its use is not limited to files. However, the way it was described is the only way it is used in this book.

called the control flow or flow of control, and statements that affect it are called control statements.†

Some kinds of compound statements can or must have more than one *clause*. The first line of each clause of a compound statement—its header in Python terminology—is at the same level of indentation as the headers of the statement's other clauses. Each header begins with a keyword and ends with a colon. The rest of the clause—its suite is a series of statements indented one level more than its header.



The term "suite" comes from Python's technical documentation. We'll generally use the more common term block instead. Also, when discussing compound statements, we frequently refer to clauses by the keywords that introduce them (for example, "a with clause").

Figure 4-1 illustrates the structure of a multi-clause compound statement. Not all compound statements are multi-clause, but every clause has a header and a suite containing at least one statement (if only pass).

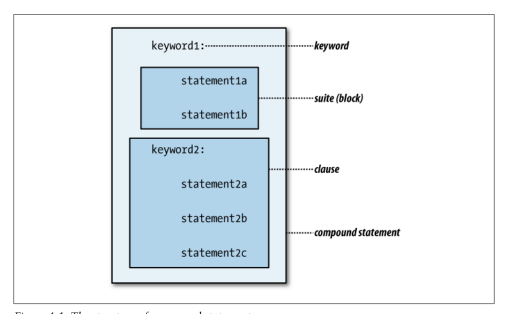


Figure 4-1. The structure of compound statements

† Strictly speaking, a function call expression (not a function definition) also affects the flow of control since it causes execution to proceed in the body of the function. Function calls can appear as separate statements or as part of expressions; either way, execution of the statements that follow or the rest of the expression is suspended until the function returns. From the point of view of the code calling the function, it is a single step that does not affect the order in which statements are executed.

The statements discussed in this chapter are the scaffolding on which you will build your programs. Without them you are limited to the sort of simple computations shown in the examples of the previous chapters. The four kinds of compound statements introduced here are:

- Conditionals
- Loops
- Iterations
- Exception handlers



Starting later in this chapter, some common usage patterns of Python functions, methods, and statements will be presented as abstract templates, along with examples that illustrate them. These templates are a device for demonstrating and summarizing how these constructs work while avoiding a lot of written description. They contain a mixture of Python names, self-descriptive "roles" to be replaced by real code, and occasionally some "pseudocode" that describes a part of the template in English rather than actual Python code.

The templates are in no way part of the Python language. In addition to introducing new programming constructs and techniques as you read, the templates are designed to serve as references while you work on later parts of the book and program in Python afterwards. Some of them are quite sophisticated, so it would be worth reviewing them periodically.

Conditionals

The most direct way to affect the flow of control is with a conditional statement. Conditionals in Python are compound statements beginning with if.

STATEMENT

Simple Conditional

The basic form of a conditional statement controls whether or not a block of statements will get executed.

if expression: statements

If the expression is true, the statements are executed; otherwise, they are skipped. This is like the if inside a conditional comprehension, but it's more powerful since it is followed by statements, whereas a comprehension is limited to expressions.

During the import of a module name is bound to the name of the module, but while the file is being executed name is bound to '_main_'. This gives you a way to include statements in your Python files that are executed only when the module is run or, conversely, only when it is imported. The comparison of name to 'main' would almost always be done in a conditional statement and placed at the end of the file.

A common use for this comparison is to run tests when the module is executed, but not when it is imported for use by other code. Suppose you have a function called do tests that contains lots of assignment and assertion statements that you don't want to run when the module is imported in normal use, but you do want to execute when the module is executed. While informal, this is a useful technique for testing modules you write. At the end of the file you would write:

```
if __name__ == '__main__':
    do tests()
```

There are several forms of conditional statements. The next one expresses a choice between two groups of statements and has two clauses, an if and an else.

STATEMENT

One-Alternative Conditional

In this form of conditional, the statements in the first block are executed if the expression is true; otherwise, the statements in the second block are executed.

```
if expression:
   statements1
else.
    statements2
```

A simple use of the one-alternative form of conditional is to expand the test for whether a file is being imported as opposed to being executed. We can set it up so that one thing happens when the file is imported and a different thing happens when it's executed.

This example shows only one statement in each block. There could be others, but another reason to group statements into simple functions is so you can invoke them "manually" in the interpreter during development and testing. You might run do tests, fix a few things, then run it again. These test functions are useful whether invoked automatically or manually:

```
if name == ' main ':
   do tests()
else:
   print( name , 'has been imported.')
```

The third form of conditional statement contains more than one test. Except for the if at the beginning, all the test clauses are introduced by the keyword elif.

STATEMENT

Multi-Test Conditional

This form of the statement contains an if clause and any number of elif clauses. The statement may end with an else clause, but that is not necessary.

The expressions in the if clause and each of the elif clauses are evaluated in order until one is true. Then, that clause's statements are executed and the rest of the conditional is skipped. If none of the expressions is true and there is an else clause, its statements get executed; otherwise, none of the statements are executed.

```
if expression1:
    statements1
elif expression2:
    statements2
# . . . any number of additional elif clauses
    statements
```

Python has a rich repertoire of mechanisms for controlling execution. Many kinds of maneuvers that would have been handled in older languages by conditionals—and could still be in Python—are better expressed using these other mechanisms. In particular, because they emphasize values rather than actions, conditional expressions or conditional comprehensions are often more appropriate than conditional statements.

Programming languages have many of the same properties as ordinary human languages. Criteria for clear writing are similar in both types of language. You want what vou write to be:

- Succinct
- Clear
- Accurate



It's important not to burden readers of your code (you included!) with too many details. People can pay attention to only a few things at once. Conditionals are a rather heavy-handed form of code that puts significant cognitive strain on the reader. With a little experience and experimentation, you should find that you don't often need them. There will be examples of appropriate uses in the rest of this chapter, as well as in later ones. You should observe and absorb the style they suggest.

Loops

A *loop* is a block of statements that gets executed as long as some condition is true. Loops are expressed in Python using while statements.

STATEMENT

Loop

The basic form of loop begins with the keyword while and an expression.

while expression: statements

If the expression is true, the statements are executed and the expression is evaluated again. As long as the expression is true, the statements are executed repeatedly. Once the expression is false, the statements are skipped, completing the execution of the while statement.

Note that the test may well be false the first time it is evaluated. In that case, the statements of the block won't get executed at all. If you want some code to execute once the test is false, include an else clause in your loop.

STATEMENT

Loop with Final Clause

This form of loop statement adds an else clause whose statements are executed after the expression evaluates to false.

while expression: statements1 else:xs statements2

There are two simple statements that are associated with both loops and iterations (the subject of the next section): continue and break.

STATEMENT

Loop Interruption

Occasionally it is useful to interrupt the execution of a loop's statements. The continue statement causes execution of the loop to return to the test. The break statement interrupts the entire loop's execution, causing the program to continue with the statement following the while.

The continue statement is rarely used in Python programming, but it's worth mentioning it here in case you run across it while reading someone else's Python code. The break statement is seen somewhat more often, but in most cases it is better to embed in the loop's test all the conditions that determine whether it should continue rather than using break. Furthermore, in many cases the loop is the last statement of a function, so you can just use a return statement to both end the loop and exit the function. (A return exits the function that contains it even if execution is in the middle of a conditional or loop.) Using a return instead of break is more convenient when each function does just one thing: most uses of break are intended to move past the loop to execute code that appears later in the function, and if there isn't any code later in the function a return statement inside the loop is equivalent to a break.

An error that occurs during the execution of a loop's test or one of its statements also terminates the execution of the loop. Altogether, then, there are three ways for a loop's execution to end:

Normally

The test evaluates to false.

Abnormally

An error occurs in the evaluation of the test or body of the loop.

Prematurely

The body of the loop executes a return or break statement.

When you write a loop, you must make sure that the test expression eventually becomes false or a break or return is executed. Otherwise, the program will get stuck in what is called an *infinite loop*. We'll see at the end of the chapter how to control what happens when errors occur, rather than allowing them to cause the program to exit abnormally.

Simple Loop Examples

Example 4-1 presents the simplest possible loop, along with a function that reads a line typed by the user, prints it out, and returns it.

```
Example 4-1. Echo

def echo():
    """Echo the user's input until an empty line is entered"""
    while echo1():
        pass

def echo1():
        """Prompt the user for a string, "echo" it, and return it"""
        line = input('Say something: ')
        print('You said', line)
        return line
```

The function echo1 reads a line, prints it, and returns it. The function echo contains the simplest possible while statement. It calls a function repeatedly, doing nothing (pass),

until the function returns something false. If the user just presses Return, echo1 will print and return an empty string. Since empty strings are false, when the while gets an empty string back from echo1 it stops. A slight variation, shown in Example 4-2, is to compare the result returned from echo1 to some specified value that signals the end of the conversation.

```
Example 4-2. Polite echo
def polite echo():
     """Echo the user's input until it equals 'bye'"""
    while echo1() != 'bye':
         pass
```

Of course, the bodies of loops are rarely so trivial. What allows the loop in these examples to contain nothing but a pass is that echo1 is called both to perform an action and to return True or False. This example uses such trivial loop bodies only to illustrate the structure of the while statement.

Initialization of Loop Values

Example 4-3 shows a more typical loop. It records the user's responses, and when the user types 'bye' the function returns a record of the input it received. The important thing here is that it's not enough to use echo1's result as a test. The function also needs to add it to a list it is building. That list is returned from the function after the loop exits.

```
Example 4-3. Recording echo
```

```
def recording echo():
    """Echo the user's input until it equals 'bye', then return a list of all the inputs received"""
    lst = []
    entry = echo1()
    while entry != 'bye':
         lst.append(entry)
         entry = echo1()
    return 1st
```

In this example, echo1 is called in two places: once to get the first response and then each time around the loop. Normally it is better not to repeat a piece of code in two places, even if they are so close together. It's easy to forget to change one when you change the other or to make incompatible changes, and changing the same thing in multiple places is tedious and error-prone. Unfortunately, the kind of repetition shown in this example is often difficult to avoid when combining input—whether from the user or from a file—with while loops.

TEMPLATE

Function with a Simple Loop

As small as Example 4-3 is, it demonstrates all the usual parts of functions containing a while statement.

```
def fn():
    initialize values
    while test values:
        use values
        change values
        # repeat
    return result
```

Example 4-4 shows the same function as Example 4-3, but with comments added to emphasize the way the code uses a simple loop.

Example 4-4. Commented recording echo function

```
def recording_echo():
    # initialize entry and lst
    lst = []
    # get the first input
    entry = echo1()

# test entry
    while entry != 'bye':

    # use entry
    lst.append(entry)

# change entry
    entry = echo1()

# repeat

# return result
return lst
```

All parts of this template are optional except for the line beginning with while. Typically, one or more of the values assigned in the initialization portion are used in the loop test and changed inside the loop. In recording_echo the value of entry is initialized, tested, used, and changed; lst is initialized, used, and changed, but it is not part of the loop's test.

Looping Forever

Sometimes you just want your code to repeat something until it executes a **return** statement. In that case there's no need to actually test a value. Since while statements require a test, we use True, which is, of course, always true. This may seem a bit odd, but there are times when something like this is appropriate. It is often called "looping forever." Of course, in reality the program won't run "forever," but it might run forever as far as *it* is concerned—that is, until something external causes it to stop. Such programs are found frequently in operating system and server software.

TEMPLATE

Loop Forever

A conditional's loop expression can be as simple as a single true value, causing it to loop until an external event stops the program.

```
initialize values
while True:
    change values
    if test values:
        return
    use values
    # repeat
return result
```

Example 4-5 shows a rewrite of Example 4-3 using the Loop Forever template. Typical loops usually get the next value at the end of the loop, but in this kind, the next value is obtained at the beginning of the loop.

Example 4-5. Recording echo using "loop forever"

```
def recording echo with conditional():
     """Echo the user's input until it equals 'bye', then return a list of all the inputs received"""
    # no need to initialize a value to be tested since nothing is tested!
    while True:
         entry = echo1()
         if entry == 'bye':
             return sea
         seq.append(entry)
```

Loops over generators are always effectively "forever" in that there's no way to know how many items the generator will produce. The program must call next over and over again until the generator is exhausted. We saw in Chapter 3 (in "Generators" on page 78) that the generator argument of next can be followed by a value to return when the generator is exhausted. A "forever" loop can be written to use this feature in a function that combines all of the generated amino acid abbreviations into a string. Example 4-6 repeats the definition of the generator function and shows the definition of a new function that uses it.

Example 4-6. Looping over a generator of amino acid symbols

```
def aa generator(rnaseq):
     """Return a generator object that produces an amino acid by translating
    the next three characters of rnaseq each time next is called on it"""
    return (translate RNA codon(rnaseq[n:n+3])
             for n in range(0, len(rnaseq), 3))
def translate(rnaseq):
    """Translate rnaseq into amino acid symbols"""
    gen = aa generator(rnaseq)
```

```
seq = ''
aa = next(gen, None)
while aa:
    seq += aa
    aa = next(gen, None)
return seq
```

Loops with Guard Conditions

Loops are often used to search for a value that meets the test condition when there is no guarantee that one does. In such situations it is not enough to just test each value—when there are no more values, the test would be repeated indefinitely. A second conditional expression must be added to detect that no more values remain to be tested.

TEMPLATE

Search Loop

In a loop that searches for a value that meets a particular test, the loop condition takes the form:

```
not end and not test
```

The loop stops when either there are no more values (*end* is true) or a value passes the test. If the function is intended to return a value it must test *end* again after the loop, returning None if it is true and the value that met the test if it is false.

```
initialize values
while not at-end and not at-target:
    use current values
    get new values
    # repeat
return success-result if test else None
```

Loops like these are used when there are two separate reasons for them to end: either there are no more values to use—at-end—or some kind of special value has been encountered, detected by at-target. If there are no more values to consider, evaluating at-target would be meaningless, or, as is often the case, would cause an error. The and operator is used to "protect" the second part of the test so that it is evaluated only if the first is true. This is sometimes called a guard condition.

When a loop can end for more than one reason, the statements after the while will need to distinguish the different cases. The simplest and most common case is to return one value if the loop ended because *at-end* became true and a different value if the loop ended because *at-target* became true.

Two-condition loops like this occur frequently in code that reads from streams such as files, terminal input, network connections, and so on. That's because the code cannot know when it has reached the end of the data until it tries to read past it. Before the result of a read can be used, the code must check that something was actually read.

Because readline returns '\n' when it reads a blank line but returns '' at the end of a file, it is sufficient to check to see that it returned a nonempty string. Repeated calls to readline at the end of the file will continue to return empty strings, so if its return value is not tested the loop will never terminate.

Example 4-7 shows a function that reads the first sequence from a FASTA file. Each time it reads a line it must check first that the line is not empty, indicating that the end of file has been reached, and if not, that the line does not begin with '>', indicating the beginning of the next sequence.

Example 4-7. Checking for a result after every read

```
def read sequence(filename):
    """Given the name of a FASTA file named filename, read and return
    its first sequence, ignoring the sequence's description"""
    seq = ''
    with open(filename) as file:
        line = file.readline()
        while line and line[0] == '>':
            line = file.readline()
        while line and line[0] != '>':
                                                      # must check for end of file
            sea += line
            line = file.readline()
    return seq
```



Although files can often be treated as collections of lines using comprehensions or readlines, in some situations it is more appropriate to loop using readline. This is especially true when several related functions all read from the same stream.

The bare outline of code that loops over the lines of a file, doing something to each, is shown in the next template.

TEMPLATE

Looping Over Lines of a File

When several related functions each loop over lines of a file while testing for an end condition that terminates their processing, each must check whether the end of the file has been reached before performing its test. Each function's loop would have the form:

```
line = file.readline()
while line and not test(line):
    do something with line
    line = file.readline()
```

Iterations

Collections contain objects; that's more or less *all* they do. Some built-in functions—min, max, any, and all—work for any type of collection. The operators in and not in accept any type of collection as their second operand. These functions and operators have something very important in common: they are based on doing something with each element of the collection.‡ Since any element of a collection could be its minimum or maximum value, min and max must consider all the elements. The operators in and not in and the functions any and all can stop as soon as they find an element that meets a certain condition, but if none do they too will end up considering every element of the collection.

Doing something to each element of a collection is called *iteration*. We've actually already seen a form of iteration—comprehensions. Comprehensions "do something" to every element of a collection, collecting the results of that "something" into a set, list, or dictionary. Comprehensions with multiple for clauses perform nested iterations. Comprehensions with one or more if clauses perform conditional iteration: unless an element passes all the if tests, the "something" will not be performed for that element and no result will be added.

The code in this book uses comprehensions much more aggressively than many Python programmers do. You should get comfortable using them, because in applicable situations they say just what they mean and say it concisely. Their syntax emphasizes the actions and tests performed on the elements. They produce collection objects, so the result of a comprehension can be used in another expression, function call, return statement, etc. Comprehensions help reduce the littering of code with assignment statements and the names they bind.

The question, then, is: what kinds of collection manipulations do *not* fit the mold of Python's comprehensions? From the point of view of Python's language constructs, the answer is that actions performed on each element of a collection sometimes must be expressed using statements, and comprehensions allow only expressions. Comprehensions also can't stop before the end of the collection has been reached, as when a target value has been located. For these and other reasons Python provides the for statement to perform general-purpose iteration over collections.

Iteration Statements

Iteration statements all begin with the keyword for. This section shows many ways for statements can be used and "templates" that summarize the most important.

[‡] The len function is a bit different: while it could be implemented by counting the elements one at a time, most types implement length more directly.

STATEMENT

Iteration

Python's for statement expresses iteration succinctly:

for item in collection: do something with item

You will use for statements often, since so much of your programming will use collections. The for statement makes its purpose very clear. It is easy to read and write and minimizes the opportunities for making mistakes. Most importantly, it works for collections that aren't sequences, such as sets and dictionaries. As a matter of fact, the for statement isn't even restricted to collections: it works with objects of a broader range of types that together are categorized as iterables. (For instance, we are treating file objects as collections (streams), but technically they are another kind of iterable.)



The continue and break statements introduced in the section on loops work for iterations too.

TEMPLATE

File Iteration

Doing something with each line of a file object is an especially useful application of the for statement.

with open(filename) as file: for line in file: do something with line

By default, a dictionary iteration uses the dictionary's keys. If you want the iteration to use its values, call the values method explicitly. To iterate with both keys and values at the same time, call the items method. Typically, when using both keys and values you would unpack the result of items and assign a name to each, as shown at the end of the following template.

TEMPLATE

Dictionary Iteration

Iteration can be performed with a dictionary's keys, values, or key/value pairs.

for key in dictionary.keys(): do something with key

```
for value in dictionary.values():
    do something with value
```

While an iteration that uses a dictionary's items could begin with for item in, it is usually preferable to use tuple unpacking to name the key and value in each item tuple.

```
for key, value in dictionary.items():
   do something with key and value
```

The previous chapter pointed out that if you need a dictionary's keys, values, or items as a list you can call list on the result of the corresponding method. This isn't necessary in for statements—the results of the dictionary methods can be used directly. keys, values, and items each return an iterable of a different type—dict keys, dict values, and dict items, respectively—but this difference almost never matters, since the results of calls to these methods are most frequently used in for statements and as arguments to list.

Sometimes it is useful to generate a sequence of integers along with the values over which a for statement iterates. The function enumerate(iterable) generates tuples of the form (n, value), with n starting at 0 and incremented with each value taken from the iterable. It is rarely used anywhere but in a for statement.

TEMPLATE

Numbering Iterations

Use the function enumerate and tuple unpacking to generate numerical keys in parallel with the values in an iterable.

```
for n, value in enumerate(iterable):
   do something with n and value
```

A common use for enumerate is to print out the elements of a collection along with a sequence of corresponding numbers. The "do something" line of the template becomes a call to print like the following:

```
print(n, value, sep='\t')
```

Kinds of Iterations

Most iterations conform to one of a small number of patterns. Templates for these patterns and examples of their use occupy most of the rest of this chapter. Many of these iteration patterns have correlates that use loops. For instance, Example 4-3 is just like the Collect template described shortly. In fact, anything an iteration can do can be done with loops. In Python programming, however, loops are used primarily to deal with external events and to process files by methods other than reading them line by line.



Iteration should always be preferred over looping. Iterations are a clearer and more concise way to express computations that use the elements of collections, including streams. Writing an iteration is less error-prone than writing an equivalent loop, because there are more details to "get right" in coding a loop than in an iteration.

Do

Often, you just want to do something to every element of a collection. Sometimes that means calling a function and ignoring its results, and sometimes it means using the element in one or more statements (since statements don't have results, there's nothing to ignore).

TEMPLATE

D٥

The simplest kind of iteration just does something for each element of a collection.

for item in collection: do something with item

A very useful function to have is one that prints every element of a collection. When something you type into the interpreter returns a long collection, the output is usually difficult to read. Using pprint.pprint helps, but for simple situations the solution demonstrated in Example 4-8 suffices. Both pprint and this definition can be used in other code too, of course.

Example 4-8. Doing something to (print) each element

```
def print collection(collection):
   for item in collection:
        print(item)
   print()
```

Actually, even this action could be expressed as a comprehension:

```
[print(item) for item in collection]
```

Since print returns None, what you'd get with that comprehension is a list containing one None value for each item in the collection. It's unlikely that you'd be printing a list large enough for it to matter whether you constructed another one, but in another situation you might call some other no-result function for a very large collection. However, that would be silly and inefficient. What you could do in that case is use a set, instead of list, comprehension:

{print(item)for item in collection}

That way the result would simply be {None}. Really, though, the use of a comprehension instead of a Do iteration is not a serious suggestion, just an illustration of the close connection between comprehensions and iterations.

We can create a generalized "do" function by passing the "something" as a functional argument, as shown in Example 4-9.

Example 4-9. A generalized do function

```
def do(collection, fn):
   for item in collection:
       fn(item)
```

The function argument could be a named function. For instance, we could use do to redefine print collection from Example 4-8, as shown in Example 4-10.

Example 4-10. Redefining print_collection using a generalized do

```
def print collection(collection):
   do(collection, print)
```

This passes a named function as the argument to do. For more ad hoc uses we could pass a lambda expression, as in Example 4-11.

```
Example 4-11. Passing a lambda expression to do
do(collection, lambda elt: print('\t', elt, sep=''))
```

The way to express a fixed number of repetitions of a block of code is to iterate over a range, as shown in the following template.

TEMPLATE

Repeat

To repeat a block of statements n times, iterate over range(n).

```
for count in range(n):
   statements
```

Frequently, count would not even be used in the body of the iteration.

Collect

Iterations often collect the results of the "something" that gets done for each element. That means creating a new list, dictionary, or set for the purpose and adding the results to it as they are computed.

TEMPLATE

Collect

A Collect iteration starts with an empty collection and uses a method or operator appropriate to its type to add something to it for each iteration.

```
result = []
for item in collection:
   statements using item
   result.append(expression based on the statements)
return result
```

Most situations in which iteration would be used to collect results would be better expressed as comprehensions. Sometimes, though, it can be tricky to program around the limitation that comprehensions cannot contain statements. In these cases, a Collect iteration may be more straightforward. Perhaps the most common reason to use a Collect iteration in place of a comprehension or loop is when one or more names are assigned and used as part of the computation. Even in those cases, it's usually better to extract that part of the function definition and make it a separate function, after which a call to that function can be used inside a comprehension instead of an iteration.

Example 4-12 shows a rewrite of the functions for reading entries from FASTA files in Chapter 3. In the earlier versions, all the entries were read from the file and then put through a number of transformations. This version, an example of the Collect iteration template, reads each item and performs all the necessary transformations on it before adding it to the collection. For convenience, this example also repeats the most succinct and complete comprehension-based definition.

While more succinct, and therefore usually more appropriate for most situations, the comprehension-based version creates several complete lists as it transforms the items. Thus, with a very large FASTA file the comprehension-based version might take a lot more time or memory to execute. After the comprehension-based version is yet another, this one using a loop instead of an iteration. You can see that it is essentially the same, except that it has extra code to read the lines and check for the end of the file.

Example 4-12. Reading FASTA entries with a Collect iteration

```
def read FASTA iteration(filename):
   sequences = []
   descr = None
   with open(filename) as file:
        for line in file:
            if line[0] == '>':
                if descr:
                                                          # have we found one yet?
                    sequences.append((descr, seq))
                descr = line[1:-1].split('|')
                seq = ''
                                                          # start a new sequence
            else:
                seq += line[:-1]
```

```
sequences.append((descr, seq))
                                                          # add the last one found
    return sequences
def read FASTA(filename):
    with open(filename) as file:
        return [(part[0].split('|'),
                 part[2].replace('\n', ''))
                for part in
                [entry.partition('\n')
                 for entry in file.read().split('>')[1:]]]
def read FASTA loop(filename):
    sequences = []
    descr = None
    with open(filename) as file:
        line = file.readline()[:-1]
                                                          # always trim newline
        while line:
            if line[0] == '>':
                if descr:
                                                          # any sequence found yet?
                     sequences.append((descr, seq))
                descr = line[1:].split('|')
                seq = ''
                                                          # start a new sequence
            else:
                seq += line
            line = file.readline()[:-1]
        sequences.append((descr, seq))
                                                          # easy to forget!
    return sequences
```

Combine

Sometimes we want to perform an operation on all of the elements of a collection to yield a single value. An important feature of this kind of iteration is that it must begin with an initial value. Python has a built-in sum function but no built-in product; Example 4-13 defines one.

```
Example 4-13. A definition of product
```

As simple as this definition is, there is no reasonable way to define it just using a comprehension. A comprehension always creates a collection—a set, list, or dictionary—and what is needed here is a single value. This is called a Combine (or, more technically, a "Reduce"§) because it starts with a collection and ends up with a single value.

[§] The term "reduce" comes from the mathematical idea that a one-dimensional collection is *reduced* to a "zero"-dimensional "scalar" value.

TEMPLATE

Combine

The general outline for a Combine iteration is as follows, with \cdot representing a binary operation and fn a two-parameter function. The *identity value* is the value i for which $i \cdot v == v$ or fn(i,v) == v. Typical examples would be 0 for addition and 1 for multiplication. When collections are being combined, the identity element is an empty collection.

Only one of the three forms shown inside the for statement would be used; they are shown here together for convenience.

```
result = identity-value
for item in collection:
    result = result · item
                                      # One of these
                                      # three forms
    result ⋅= item
   result = fn(result, item)
                                      # is used.
return result
```

For another example, let's find the longest sequence in a FASTA file. We'll assume we have a function called read FASTA, like one of the implementations shown in Chapter 3. Example 4-13 used a binary operation to combine each element with the previous result. Example 4-14 uses a two-valued function instead, but the idea is the same. The inclusion of an assignment statement inside the loop is an indication that the code is doing something that cannot be done with a comprehension.

Example 4-14. Combine: identifying the longest FASTA sequence

```
def longest sequence(filename):
   longest seq = ''
   for info, seg in read FASTA(filename):
        longest seq = max(longest seq, seq, key=len)
   return longest seq
```

A special highly reduced form of Combine is Count, where all the iteration does is count the number of elements. It would be used to count the elements in an iterable that doesn't support length. This template applies particularly to generators: for a generator that produces a large number of items, this is far more efficient than converting it to a list and then getting the length of the list.

TEMPLATE

Count

A Count iteration "combines" the value 1 for each element of the iteration.

```
count = 0
for item in iterable:
   count += 1
return count
```

One of the most important and frequently occurring kinds of actions on iterables that cannot be expressed as a comprehension is one in which the result of doing something to each element is itself a collection (a list, usually), and the final result is a combination of those results. An ordinary Combine operation "reduces" a collection to a value; a Collection Combine reduces a *collection of collections* to a *single collection*. (In the template presented here the reduction is done step by step, but it could also be done by first assembling the entire collection of collections and then reducing them to a single collection.)

TEMPLATE

Collection Combine

In this variation on Combine, an action is performed on each element of a collection that produces a collection as a result, but instead of returning a collection of the results, the iteration combines the results into (reduces the results to) a single collection.

```
result = []
for item in collection:
    result += fn(item)
    # merge result with previous results
return result
```

Example 4-15 shows an example in which "GenInfo" IDs are extracted from each of several files, and a single list of all the IDs found is returned.

Example 4-15. Collection Combine: sequence IDs from multiple files

```
def extract gi id(description):
    """Given a FASTA file description line, return its GenInfo ID if it has one"""
        if line[0] != '>':
        return None
    fields = description[1:].split('|')
    if 'gi' not in fields:
        return None
    return fields[1 + fields.index('gi')]
def get gi ids(filename):
     """Return a list of the GenInfo IDs of all sequences found in the file named filename"""
    with open(filename) as file:
        return [extract gi id(line) for line in file if line[0] == '>']
def get gi ids from files(filenames):
     ""Return a list of the GenInfo IDs of all sequences found in the
    files whose names are contained in the collection filenames"""
    idlst = []
    for filename in filenames:
        idlst += get gi ids(filename)
    return idlst
```

Search

Another common use of iterations is to search for an element that passes some kind of test. This is not the same as a combine iteration—the result of a combination is a property of all the elements of the collection, whereas a search iteration is much like a search loop. Searching takes many forms, not all of them iterations, but the one thing you'll just about always see is a return statement that exits the function as soon as a matching element has been found. If the end of the function is reached without finding a matching element the function can end without explicitly returning a value, since it returns None by default.

TEMPLATE

Search

Search is a simple variation on Do:

for item in collection: if test item: return item

Suppose we have an enormous FASTA file and we need to extract from it a sequence with a specific GenBank ID. We don't want to read every sequence from the file, because that could take much more time and memory than necessary. Instead, we want to read one entry at a time until we locate the target. This is a typical search. It's also something that comprehensions can't do: since they can't incorporate statements, there's no straightforward way for them to stop the iteration early.

As usual, we'll build this out of several small functions. We'll define four functions. The first is the "top-level" function; it calls the second, and the second calls the third and fourth. Here's an outline showing the functions called by the top-level function:

```
search_FASTA_file_by_gi_id(id, filename)
    FASTA search by gi id(id, fil)
        extract gi id(line)
       read FASTA sequence(fil)
```

This opens the file and calls FASTA search by gi id to do the real work. That function searches through the lines of the file looking for those beginning with a '>'. Each time it finds one it calls get gi id to get the GenInfo ID from the line, if there is one. Then it compares the extracted ID to the one it is looking for. If there's a match, it calls read FASTA sequence and returns. If not, it continues looking for the next FASTA description line. In turn, read FASTA sequence reads and joins lines until it runs across a description line, at which point it returns its result. Example 4-16 shows the definition of the top-level function.

"Top-level" functions should almost always be very simple. They are entry points into the capabilities the other function definitions provide. Essentially, what they do is prepare the information received through their parameters for handling by the functions that do the actual work.

Example 4-16. Extracting a sequence by ID from a large FASTA file

Each of the other functions can be implemented in two ways. Both FASTA_search_by_gi_id and read_FASTA_sequence can be implemented using a loop or iteration. The simple function get_gi_id can be implemented with a conditional expression or a conditional statement. Table 4-1 shows both implementations for FASTA_search_by_gi_id.

Table 4-1. Two definitions of FASTA_search_by_gi_id

```
Iteration
                                                   Loop
                         def FASTA search by gi id(id, file):
for line in file:
                                                   line = file.readline()
    if (line[0] == '>' and
                                                   while (line and
                                                          not (line[0] == '>' and
        str(id) == get_gi_id(line)):
           return \
                                                                (str(id) ==
                                                                   get_gi_id(line)))):
               read FASTA sequence(file)
                                                                line = file.readline()
                                                   return (line and
                                                          read_FASTA_sequence(fil)
```

The iterative implementation of FASTA_search_by_gi_id treats the file as a collection of lines. It tests each line to see if it is the one that contains the ID that is its target. When it finds the line it's seeking, it does something slightly different than what the template suggests: instead of returning the line—the item found—it goes ahead and reads the sequence that follows it.



The templates in this book are not meant to restrict your code to specific forms: they are frameworks for you to build on, and you can vary the details as appropriate.

The next function—read_FASTA_sequence—shows another variation of the search template. It too iterates over lines in the file—though not all of them, since it is called after FASTA_search_by_gi_id has already read many lines. Another way it varies from the template is that it accumulates a string while looking for a line that begins with a '>'. When it finds one, it returns the accumulated string. Its definition is shown in Table 4-2, and the definition of get_gi_id is shown in Table 4-3.

Table 4-2. Two definitions of read_FASTA_sequence

```
Iteration
                                                    Loop
                          read FASTA sequence(file):
                              seq =
for line in file:
                                                    line = file.readline()
                                                    while line and line[0] != '>':
    if not line or line[0] == '>':
                                                         seq += line[:-1]
        return seq
   seq += line[:-1]
                                                        line = file.readline()
                                                    return seq
```

Table 4-3. Two definitions of get_gi_id

```
Conditional statement
                                                     Conditional expression
                          def get gi id(description):
                               fields = description[1:].split('|')
if fields and 'gi' in fields:
                                                     return (fields and 'gi' in fields and
   return fields[(1 +
                                                             fields[1+fields.index('gi')])
                   fields.index('gi')]
```

A special case of search iteration is where the result returned is interpreted as a Boolean rather than the item found. Some search iterations return False when a match is found, while others return True. The exact form of the function definition depends on which of those cases it implements. If finding a match to the search criteria means the function should return False, then the last statement of the function would have to return True to show that all the items had been processed without finding a match. On the other hand, if the function is meant to return True when it finds a match it is usually not necessary to have a return at the end, since None will be returned by default, and None is interpreted as false in logical expressions. (Occasionally, however, you really need the function to return a Boolean value, in which case you would end the function by returning False.) Here are two functions that demonstrate the difference:

```
def rna sequence is valid(seq):
    for base in seq:
        if base not in 'UCAGucag':
            return False
    return True
def dna sequence contains N(seq):
    for base in seq:
    if base == 'N':
        return True
```

Filter

Filtering is similar to searching, but instead of returning a result the first time a match is found, it does something with each element for which the match was successful. Filtering doesn't stand on its own—it's a modification to one of the other kinds of iterations. This section presents templates for some filter iterations. Each just adds a

conditional to one of the other templates. The condition is shown simply as **test** item, but in practice that test could be complex. There might even be a few initialization statements before the conditional.

TEMPLATE

Filtered Do

A Filtered Do performs an action for each item that meets a specified condition.

```
for item in collection:
   if test item:
       statements using item
```

An obvious example of a Filtered Do is printing the header lines from a FASTA file. Example 4-17 shows how this would be implemented.

Example 4-17. Printing the header lines from a FASTA file

As with Collect iterations in general, simple situations can be handled with comprehensions, while iterations can handle the more complex situations in which statements are all but unavoidable. For example, extracting and manipulating items from a file can often be handled by comprehensions, but if the number of items is large, each manipulation will create an unnecessarily large collection. Rather than collecting all the items and performing a sequence of operations on that collection, we can turn this inside out, performing the operations on one item and collecting only the result.

TEMPLATE

Filtered Collect

Here, the values for which test is true are collected one at a time.

```
result = []
for item in collection:
    if test item:
        statements using item
        result.append(expression based on the statements)
return result
```

In many cases, once a line passes the test the function should not return immediately. Instead, it should continue to read lines, concatenating or collecting them, until the next time the test is true. An example would be with FASTA-formatted files, where a function might look for all sequence descriptions that contain a certain string, then

read all the lines of the sequences that follow them. What's tricky about this is that the test applies only to the lines beginning with '>'. The lines of a sequence do not provide any information to indicate whether they should be included or not.

Really what we have here are two tests: there's a preliminary test that determines whether the primary test should be performed. Neither applies to the lines that follow a description line in the FASTA file, though. To solve this problem, we add a flag to govern the iteration and set it by performing the primary test whenever the preliminary test is true. Example 4-18 shows a function that returns the sequence strings for all sequences whose descriptions contain the argument string.

Example 4-18. Extracting sequences with matching descriptions

```
def extract matching sequences(filename, string):
    """From a FASTA file named filename, extract all sequences whose descriptions contain string"""
    sequences = []
    sea = ''
    with open(filename) as file:
        for line in file:
             if line[0] == '>':
                 if seq:
                                                                # not first time through
                      sequences.append(seq)
                                # next sequence detected
                 includeflag = string in line
                                                                # flag for later iterations
             else:
                 if includeflag:
                     seq += line[:-1]
        if seq:
                                                                # last sequence in file is included
             sequences.append(seq)
    return sequences
```

The generalization of this code is shown in the following template.

TEMPLATE

Filtered Collect of Groups of Lines

The details of code implementing this template vary more than the template implies. Its overall job is to look for a line that meets both a preliminary test and a primary test. It collects or concatenates lines when the primary test is true until the next line for which the preliminary test is true. This requires a flag that keeps track of the result of the primary test while subsequent lines are read.

```
lines = []
with open(inputfilename) as file:
    for line in file:
        if preliminary-test:
            flag = primary-test(line)
            lines.append(line)
                                                            # or concatenate, etc.
return lines
```

A Filtered Combine is just like a regular Combine, except only elements that pass the test are used in the combining expression.

TEMPLATE

Filtered Combine

As with a regular Combine, only one of the three forms shown inside the condition is used for a given piece of code implementing this template.

Example 4-13 showed a definition for product. Suppose the collection passed to product contained nonnumerical elements. You might want the product function to skip nonnumerical values instead of converting string representations of numbers to numbers.

All that's needed to skip nonnumerical values is a test that checks whether the element is an integer or float and ignores it if it is not. The function <code>isinstance</code> was described briefly in Chapter 1; we'll use that here to check for numbers. Example 4-19 shows this new definition for product.

Example 4-19. Filtered Combine: another definition of product

What we've done here is replace the template's *test* with a call to <code>is_number</code> to perform the test. Suppose we needed different tests at different times while computing the product—we might want to ignore zeros or negative numbers, or we might want to start at a different initial value (e.g., 1 if computing the product of only integers). We might even have different actions to perform each time around the iteration. We can

^{||} Spreadsheet applications, for example, typically skip nonnumbers when performing numeric operations like "sum" on a row or column, rather than producing an error.

implement many of these templates as function definitions whose details are specified by parameters. Example 4-20 shows a completely general combine function.

Example 4-20. Generalized combine function

```
def combine(coll, initval, action, filter=None):
     """Starting at initval, perform action on each element of coll, finally returning the result. If
    filter is not None, only include elements for which filter(element) is true. action is a function
    of two arguments--the interim result and the element--which returns a new interim result."""
    result = initval
    for elt in coll:
         if not filter or filter(elt):
            result = action(result, elt)
    return result
```

To add all the integers in a collection, we just have to call combine with the right arguments:

```
combine(coll
        lambda result, elt: result + elt,
        lambda elt: isinstance(elt, int)
```

TEMPLATE

Filtered Count

An important specific variation of Filtered Combine is Filtered Count. This is useful for collections even if they support the len function, because len cannot include certain items.

```
count = 0
for item in iterable:
    if test item:
        count += 1
return count
```

Nested iterations

One iteration often uses another. Example 4-21 shows a simple case—listing all the sequence IDs in files whose names are in a collection.

Example 4-21. A nested iteration

```
def list sequences in files(filelist):
    ""For each file whose name is contained in filelist,
    list the description of each sequence it contains"""
    for filename in filelist:
        print(filename)
        with open(filename) as file:
             for line in file:
                 if line[0] == '>':
                      print('\t', line[1:-1])
```

Nesting is not a question of physical containment of one piece of code inside another. Following the earlier recommendation to write short, single-purpose functions, Example 4-22 divides the previous function, placing one iteration in each. This is still a nested iteration, because the first function calls the second each time around the for, and the second has its own for statement.

Example 4-22. A two-function nested iteration

```
def list_sequences_in_files(filelist):
    """For each file whose name is contained in filelist,
    list the description of each sequence it contains"""
    for filename in filelist:
        print(filename)
        with open(filename) as file:
            list_sequences_in_file(file)

def list_sequences_in_file(file)
    for line in file:
        if line[0] == '>':
            print('\t', line[1:-1])
```

These examples do present nested iterations, but they don't show what's special about this kind of code. Many functions that iterate call other functions that also iterate. They in turn might call still other functions that iterate. Nested iterations are more significant when their "do something" parts involve doing something with a value from the outer iteration and a value from the inner iteration together.

TEMPLATE

Nested Iteration

The general form of a nested iteration is as follows, keeping in mind that the inner iteration might actually be defined as a separate function.

```
for outer in outer_collection:
    for inner in inner_collection:
        do something with inner and outer
```

Perhaps a batch of samples is to be submitted for sequencing with each of a set of primers:

```
for seq in sequences:
    for primer in primers:
        submit(seq, primer)
```

This submits a sequence and a primer for every combination of a sequence from sequences and a primer from primers. In this case it doesn't matter which iteration is the outer and which is the inner, although if they were switched the sequence/primer pairs would be submitted in a different order.

Three-level iterations are occasionally useful—especially in bioinformatics programming, because codons consist of three bases. Example 4-23 shows a concise three-level iteration that prints out a simple form of the DNA codon table.

Example 4-23. Printing the codon table

```
def print codon table():
     ""Print the DNA codon table in a nice, but simple, arrangement"""
    for base1 in DNA bases:
                                                         # horizontal section (or "group")
        for base3 in DNA bases:
                                                         # line (or "row")
            for base2 in DNA bases:
                                                         # vertical section (or "column")
                # the base2 loop is inside the base3 loop!
                print(base1+base2+base3,
                       translate DNA codon(base1+base2+base3),
                       end=' ')
            print()
        print()
>>> print codon table()
TTT Phe TCT Ser TAT Tyr TGT Cys
TTC Phe TCC Ser TAC Tyr TGC Cys
TTA Leu TCA Ser TAA --- TGA ---
TTG Leu TCG Ser TAG --- TGG Trp
CTT Leu CCT Pro CAT His CGT Arg
CTC Leu CCC Pro CAC His CGC Arg
CTA Leu CCA Pro CAA Gln CGA Arg
CTG Leu CCG Pro CAG Gln CGG Arg
ATT Ile ACT Thr AAT Asn AGT Ser
ATC Ile ACC Thr AAC Asn AGC Ser
ATA Ile ACA Thr AAA Lys AGA Arg
ATG Met ACG Thr AAG Lys AGG Arg
GTT Val GCT Ala GAT Asp GGT Gly
GTC Val GCC Ala GAC Asp GGC Gly
GTA Val GCA Ala GAA Glu GGA Gly
GTG Val GCG Ala GAG Glu GGG Gly
```

Recursive iterations

Trees are an important class of data structure in computation: they provide the generality needed to represent branching information. Taxonomies and filesystems are good examples. A filesystem starts at the top-level directory of, say, a hard drive. That directory contains files and other directories, and those directories in turn contain files and other directories. The whole structure consists of just directories and files.

A data structure that can contain other instances of itself is said to be recursive. The study of recursive data structures and algorithms to process them is a major subject in computer science. Trees are the basis of some important algorithms in bioinformatics too, especially in the areas of searching and indexing.

While we won't be considering such algorithms in this book, it is important to know some rudimentary techniques for tree representation and iteration. A simple while or

for statement can't by itself follow all the branches of a tree. When it follows one branch, it may encounter further branches, and at each juncture it can follow only one at a time. It can only move on to the next branch after it's fully explored everything on the first one. In the meantime, it needs someplace to record a collection of the remaining branches to be processed.

Each branch is just another tree. A function that processes a tree can call *itself* to process each of the tree's branches. What stops this from continuing forever is that eventually subtrees are reached that have no branches; these are called *leaves*. A function that calls itself—or calls another function that eventually calls it—is called a *recursive function*.

Discussions of recursion are part of many programming texts and courses. It often appears mysterious until the idea becomes familiar, which can take some time and practice. One of the advantages of recursive functions is that they can express computations more concisely than other approaches, even when recursion isn't actually necessary. Sometimes the code is so simple you can hardly figure out how it does its magic!

First, we'll look at an example of one of the ways trees are used in bioinformatics. Some very powerful algorithms used in indexing and searching genomic sequences rely on what are called *suffix trees*. These are tree structures constructed so that every path from the root to a leaf produces a subsequence that is not the prefix of any other subsequence similarly obtained. The entire string from which the tree was constructed can be recovered by traversing all paths to leaf nodes, concatenating the strings encountered along the way, and collecting the strings obtained from each path. The longest string in the resulting collection is the original string. Figure 4-2 shows an example.

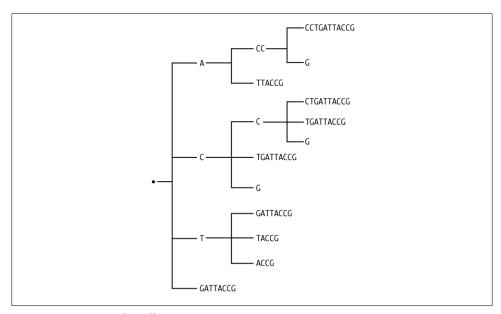


Figure 4-2. Diagram of a suffix tree

Algorithms have been developed for constructing and navigating such trees that do their work in an amount of time that is directly proportional to the length of the sequence. Normally algorithms dealing with tree-structured data require time proportional to N^2 or at best $N \log N$, where N is the length of the sequence. As N gets as large as is often required for genomic sequence searches, those quantities grow impossibly large. From this point of view the performance of suffix tree algorithms borders on the miraculous.

Our example will represent suffix trees as lists of lists of lists of... lists. The first element of each list will always be a string, and each of the rest of the elements is another list. The top level of the tree starts with an empty string. Example 4-24 shows an example hand-formatted to reflect the nested relationships.

Example 4-24. Representing a tree as a list of lists

```
],
['TTACCG']
           ['CTGATTACCG'].
           ['TGATTACCG'],
           ['G']
        ['TGATTACCG'],
        ['GATTACCG'],
        ['TACCG'],
        ['ACCG']
    ],
['GATTACCG']
 ]
```

Let's assign tree1 to this list and see what Python does with it. Example 4-25 shows an ordinary interpreter printout of the nested lists.

Example 4-25. Interpreter printing of nested lists representing a tree

```
['', ['A', ['CC', ['CCTGATTACCG'], ['G']], ['TTACCG']], ['C', ['C', ['CTGATTACCG'],
['TGATTACCG'], ['G']], ['TGATTACCG'], ['G']], ['T', ['GATTACCG'], ['TACCG'], ['ACCG']],
['GATTACCG']]
```

That output was one line, wrapped. Not very helpful. How much of an improvement does pprint.pprint offer?

```
>>> pprint.pprint(tree1)
['',
    ['A', ['CC', ['CCTGATTACCG'], ['G']], ['TTACCG']],
    ['C', ['C', ['CTGATTACCG'], ['TGATTACCG'], ['G']], ['TGATTACCG'], ['G']],
    ['T', ['GATTACCG'], ['ACCG']],
    ['GATTACCG']]
```

This is a little better, since we can at least see the top-level structure. But what we want is output that approximates the tree shown in Figure 4-2. (We won't go so far as to print symbols for lines and corners—we're just looking to reflect the overall shape of the tree represented by the data structure.) Here's the template for a recursive function to process a tree represented as just described here. (The information the tree contains could be anything, not just strings: whatever value is placed in the first position of the list representing a subtree is the value of that subtree's root node.)

TEMPLATE

Recursive Tree Iteration

This little bit of code will process all the nodes of a tree represented as a list containing a value followed by a number of subtrees. A tree with no subtrees is a list containing just a value.

```
def treewalk(tree, level=0):
    do something with tree[0] and level
    for subtree in tree[1:]:
        treewalk(node, level+1)
```

Do you find it difficult to believe that so simple a template can process a tree? Example 4-26 shows how it would be used to print our tree.

```
Example 4-26. Printing a tree
```

```
def treeprint(tree, level=0):
    print(' ' * 4 * level, tree[0], sep='')
    for node in tree[1:]:
        treeprint(node, level+1)
```

This produces the following output for the example tree. It's not as nice as the diagram; not only are there no lines, but the root of each subtree is on a line before its subtrees, rather than centered among them. Still, it's not bad for four lines of code!

```
A CC CCTGATTACCG G TTACCG C C CTGATTACCG TGATTACCG G
```

```
TGATTACCG
G
T
GATTACCG
TACCG
ACCG
GATTACCG
```

Figures 4-3, 4-4, and 4-5 illustrate the process that ensues as the function in Example 4-26 does its work with the list representing the subtree rooted at A.

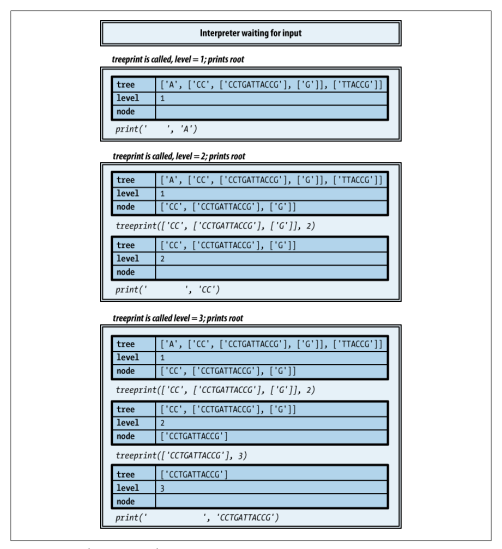


Figure 4-3. Implementation of recursion, part 1

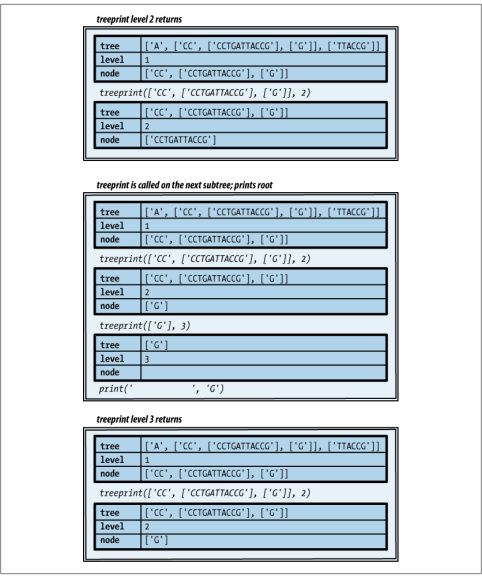


Figure 4-4. Implementation of recursion, part 2

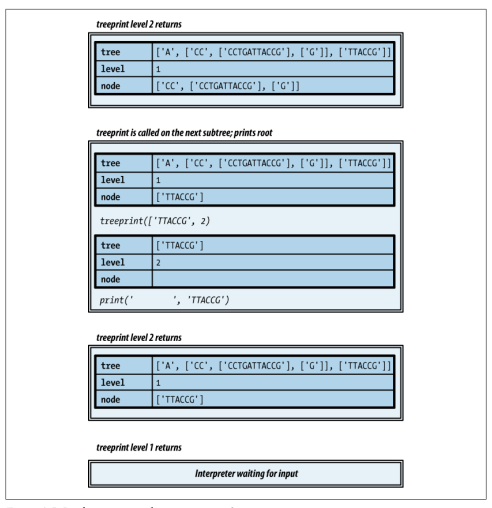


Figure 4-5. Implementation of recursion, part 3

Exception Handlers

Let's return to Example 4-15, from our discussion of collection iteration. We'll add a top-level function to drive the others and put all of the functions in one Python file called *get_gi_ids.py*. The contents of the file are shown in Example 4-27.

```
Example 4-27. Collecting GenInfo IDs of the sequences in FASTA files
```

```
def extract gi id(description):
    """Given a FASTA file description line, return its GenInfo ID if it has one"""
        if line[0] != '>':
        return None
```

```
fields = description[1:].split('|')
    if 'gi' not in fields:
        return None
    return fields[1 + fields.index('gi')]
def get gi ids(filename):
     ""Return a list of GenInfo IDs from the sequences in the FASTA file named filename"""
    with open(filename) as file:
        return [extract gi id(line) for line in file if line[0] == '>']
def get_gi_ids from files(filenames):
     ""Return a list of GenInfo IDs from the sequences in the
    FASTA files whose names are in the collection filenames"""
    idlst = []
    for filename in filenames:
        idlst += get gi ids(filename)
    return idlst
def get gi ids from user files():
    response = input("Enter FASTA file names, separated by spaces: ")
    lst = get gi ids from files(response.split()) # assuming no spaces in file names
    lst.sort()
    print(lst)
get gi ids from user files()
```

We run the program from the command line, enter a few filenames, and get the results shown in Example 4-28.

Example 4-28. Traceback from attempting to open a nonexistent file

```
% python get gi ids.py
Enter a list of FASTA filenames:
aa1.fasta aa2.fasta aa3.fasta
Traceback (most recent call last):
  File "get_gi_ids.py", line 27, in <module>
   get gi ids from user files
  File "get gi ids.py", line 23, in get gi ids from user files
   lst = get gi ids from files(files))
  File "get_gi_ids.py", line 18, in get_gi_ids_from_files
    idlst += get gi ids(filename)
  File "get_gi_ids.py", line 10, in get_gi_ids
   with open(filename) as file:
  File "/usr/local/lib/python3.1/io.py", line 278, in new
   return open(*args, **kwargs)
  File "/usr/local/lib/python3.1/io.py", line 222, in open
   closefd)
  File "/usr/local/lib/python3.1/io.py", line 619, in init
    fileio. FileIO. init (self, name, mode, closefd)
IOError: [Errno 2] No such file or directory: 'aa2.fasta'
```

Python Errors

If you've executed any Python code you have written, you have probably already seen output like that in the previous example splattered across your interpreter or shell window. Now it's time for a serious look at what this output signifies. It's important to understand more than just the message on the final line and perhaps a recognizable filename and line number or two.

Tracebacks

As its first line indicates, the preceding output shows details of pending functions. This display of information is called a traceback. There are two lines for each entry. The first shows the name of the function that was called, the path to the file in which it was defined, and the line number where its definition begins, though not in that order. (As in this case, you will often see <module> given as the module name on the first line; this indicates that the function was called from the top level of the file being run by Python or directly from the interpreter.) The second line of each entry shows the text of the line identified by the filename and line number of the first line, to save you the trouble of going to the file to read it.



Some of this will make sense to you now. Some of it won't until you have more Python knowledge and experience. As the calls descend deeper into Python's implementation, some technical details are revealed that we haven't yet explored. It's important that you resist the temptation to dismiss tracebacks as hopelessly complicated and useless. Even if you don't understand all the details, tracebacks tell you very clearly what has happened, where, and, to some extent, why.

The problem causing the traceback in this example is clear enough: the user included the file aa2.fasta in the list of files to be processed, but when get gi id went to open that file it couldn't find it. As a result, Python reported an IOError and stopped executing. It didn't even print the IDs that it had already found—it just stopped.

Runtime errors

Is this what you want your program to do? You can't prevent the user from typing the name of a nonexistent file. While you could check that each file exists before trying to open it (using methods from the os module that we'll be looking in Chapter 6), this is only one of the many things that could go wrong during the execution of your program. Maybe the file exists but you don't have read privileges for it, or it exists but is empty, and you didn't write your code to correctly handle that case. Maybe the program encounters an empty line at the end of the file and tries to extract pieces from it. Maybe the program tries to compare incompatible values in an expression such as 4 < '5'.

By now you've probably encountered ValueError, TypeError, IndexError, IOError, and perhaps a few others. Each of these errors is actually a type. Table 4-4 shows examples of common errors, the type of error instance that gets created when they occur, and examples of the messages that get printed.

Table 4-4. Common runtime errors

Example	Error class	Message
foobah	NameError	name 'foobah' is not defined
3 < '4'	TypeError	<pre>unorderable types: int() < str()</pre>
['a', 'b'] + None	TypeError	<pre>can only concatenate list (not "NoneType") to list</pre>
range()	TypeError	range expected 1 arguments, got 0
1/0	ZeroDivisionError	int division or modulo by zero
int('21', 2) ^a	ValueError	<pre>invalid literal for int() with base 2: '21'</pre>
$[1,2,3].index(4)^b$	ValueError	<pre>list.index(x): x not in list</pre>
''[1]	IndexError	string index out of range
{'a': 1}['b']	KeyError	'b'
<pre>range(4).index(5)</pre>	AttributeError	'range' object has no attribute 'index'
open('')	IOError	No such file or directory: ''
input('?')^D ^c	EOFError	
assert False	AssertionError	
assert 0, 'not 1'	AssertionError	not 1
^C^C ^d	KeyboardInterrupt	

^a The second argument of int is a base to use when reading the string that is its first argument; in this example, since base 2 was specified, only 'O' and '1' are valid in the first argument.

Even if get gi ids was written to detect nonexistent files before trying to open them, what should it do if it detects one? Should it just return None? Should it print its own error message before returning None? If it returns None, how can the function that called it know whether that was because the file didn't exist, couldn't be read, wasn't a FASTA-formatted file, or just didn't have any sequences with IDs? If each function has to report to its caller all the different problems it might have encountered, each caller will have to execute a series of conditionals checking each of those conditions before continuing with its own executions.

To manage this problem, languages provide exception handling mechanisms. These make it possible to ignore exceptions when writing most function definitions, while specifically designating, in relatively few places, what should happen when exceptions

b index is like find, but instead of returning -1 when its argument is not in the list, it raises a ValueError.

^c Typing Ctrl-D on an empty line (Ctrl-Z on Windows) ends input. Remember, though, that file.read and file.readline return empty strings if they are at the end of a file.

^d Pressing Ctrl-C twice stops whatever Python is doing and returns to the interpreter.

do occur. The term "exception" is used instead of "error" because if the program is prepared to handle a situation, it isn't really an error when it arises. It becomes an error—an unhandled exception—if the program does not detect the situation. In that case, execution stops and Python prints a traceback with a message identifying the type of error and details about the problem encountered.

Exception Handling Statements

Python's exception handling mechanism is implemented through the try statement. This looks and works much like a conditional, except that the conditions are not tests you write, but rather names of error classes.

STATEMENT **Exception Handling** The basic form of a statement that handles an exception is: try-statements except ErrorClass: except-statements

The error class is one of the error names you'll see printed out on the last line of a traceback: IOError, ValueError, and so on. When a try statement begins, it starts executing the statements in the *try-statements* block. If they complete without any errors, the rest of the try statement is skipped and execution continues at the next statement.

However, if an error of the type identified in the except clause occurs during the execution of the try block, something quite different happens: the call stack is "unwound" by removing the calls to the functions "below" the one that contains the try statement. Any of the try-statements that haven't yet executed are abandoned. Execution continues with the statements in the except clause, and then moves on to the statement that follows the entire try/except statement. Figures 4-6 and 4-7 show the difference.

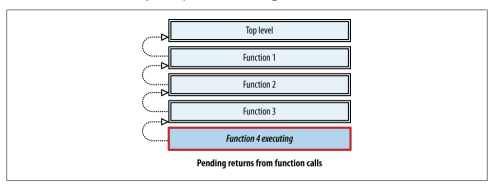


Figure 4-6. Pending returns from function calls

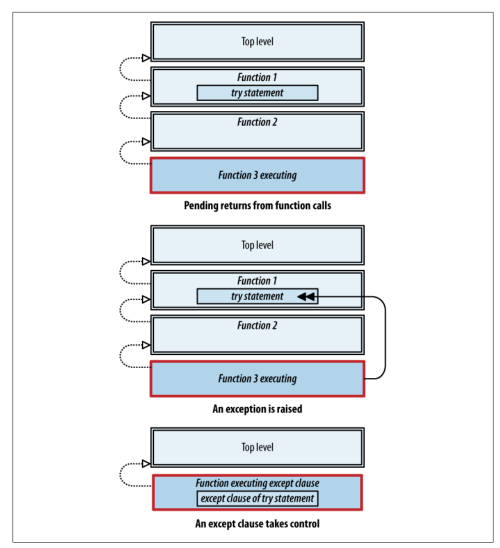


Figure 4-7. Pending returns from function calls with an exception

Optional features of exception handling statements

The try statement offers quite a few options. The difficulty here is not so much in comprehending all the details, although that does take some time. The real challenge is to develop a concrete picture of how control "flows" through function calls and Python's various kinds of statements. Then you can begin developing an understanding of the very different flow induced by exceptions.

STATEMENT

Exception Handling Options

try statement options include:

- Binding a name to the exception object caught by an except clause by following the exception class name with as and a name, allowing the statements in the clause to use details from the exception instance
- Multiple except clauses, each naming a different condition
- except clauses that specify a list of error types instead of just one
- A final except clause with no exception class to catch any exception not caught in one of the other except clauses
- A finally clause whose statements are always executed, regardless of whether an error occurs—in fact, the statements in a finally clause are executed even if the try clause or an except clause executes a return

Here's a template that shows the use of all of these features together:

```
trv:
    statements
except ErrorClass1:
    statements1
except (ErrorClass2, ErrorClass3):
    statements2
except ErrorClass4 as err:
    statements that can refer to err
    statements that are executed if an error occurs
   whose type is not in one of the above except clauses
    statements that always get executed, whether or not an error occurs
```

Now that we know how to handle errors, what changes might we want to make in our little program for finding IDs? Suppose we've decided we want the program to print an error message whenever it fails to open a file, but then continue with the next one. This is easily accomplished with one simple try statement:

```
def get gi ids(filename):
    try:
        with open(filename) as file:
            return [extract gi id(line) for line in file
                                        if line[0] == '>']
    except IOError:
        print('File', filename, 'not found or not readable.')
        return []
```

Note that the except clause returns an empty list rather than returning None or allowing the function to end without a return (which amounts to the same thing). This is because the function that calls this one will be concatenating the result with a list it is accumulating, and since None isn't a sequence it can't be added to one. (That's another

TypeError you'll often see, usually as a result of forgetting to return a value from a function.) If you've named the exception with as name, you can print(name) instead of or in addition to your own message.

Incidentally, this with statement:

```
with open('filename') as file:
        use file
is roughly the same as:
    try:
        file = open('filename')
        use file
    finally:
    file.close()
```

The finally clause guarantees that the file will be closed whether an error happens or not—and without the programmer having to remember to close it. This is a great convenience that avoids several kinds of common problems. The with statement requires only one line in addition to the statements to be executed, rather than the four lines required by the try version.

Exception handling and generator objects

An important special use of try statements is with generator objects. Each call to next with a generator object as its argument produces the generator's next value. When there are no more values, next returns the value of its optional second argument, if one is provided. If not, a StopIteration error is raised.

There are two ways to use next: either you can provide a default value and compare it to the value next returns each time, or you can omit the argument and put the call to next inside a try that has an except StopIteration clause. (An except clause with no exception class or a finally would also catch the error.)

An advantage of the exception approach is that the try statement that catches it can be several function calls back; also, you don't have to check the value returned by next each time. This is particularly useful when one function calls another that calls another, and so on. A one-argument call to next in the innermost function and a try statement in the top-level function will terminate the entire process and hand control back to the top-level function, which catches StopIteration.

Raising Exceptions

Exception raising isn't limited to library functions—your code can raise them too.

The raise statement

The raise statement is used to raise an exception and initiate exception handling.

STATEMENT

Exception Raising

The raise statement creates an instance of an exception class and begins the exception handling process.

raise exception-expression

The exception-expression can be any expression whose value is either an exception class or an instance of one. If it is an exception class, the statement simply creates an instance for you. Creating your own instance allows you to specify arguments to the new instance—typically a message providing more detail about the condition encountered. The class Exception can be used for whatever purposes you want, and it can take an arbitrary number of arguments. You can put statements like this in your code:

raise Exception('File does not appear to be in FASTA format.', filename)

The statements in any of a try statement's exception clauses can "reraise" an exception using a raise statement with no expression. In that case, the stack unwinding resumes and continues until the next try statement prepared to handle the exception is encountered.

Not only can your code raise exceptions, but you can create your own exception classes and raise instances of those. (The next chapter shows you how to create your own classes.) It's especially important for people building modules for other people to use, since code in a module has no way of knowing what code from the outside wants to do when various kinds of problems are encountered. The only reasonable thing to do is design modules to define appropriate exception classes and document them for users of the module so they know what exceptions their code should be prepared to handle.

Raising an exception to end a loop

The point was made earlier that exceptions aren't necessarily errors. You can use a combination of try and raise statements as an alternative way of ending loops. You would do this if you had written a long sequence of functions that call each other, expecting certain kinds of values in return. When something fails deep down in a sequence of calls it can be very awkward to return None or some other failure value back through the series of callers, as each of them would have to test the value(s) it got back to see whether it should continue or itself return None. A common example is repeatedly using str.find in many different functions to work through a large string.

Using exception handling, you can write code without all that distracting error reporting and checking. Exceptional situations can be handled by raising an error. The first function called can have a "while-true" loop inside a try statement. Whenever some function determines that nothing remains to process, it can throw an exception. A good exception class for this purpose is **StopIteration**, which is used in the implementation of generators, while-as statements, and other mechanisms we've seen:

```
try:
    while(True):
        begin complicated multi-function input processing
except StopIteration:
    pass
... many definitions of functions that call each other; ...
... wherever one detects the end of input, it does: ...
        raise StopIteration
```

Extended Examples

This section presents some extended examples that make use of the constructs described earlier in the chapter.

Extracting Information from an HTML File

Our first example in this section is based on the technique just discussed of raising an exception to end the processing of some text. Consider how you would go about extracting information from a complex HTML page. For example, go to NCBI's Entrez Gene site (http://www.ncbi.nlm.nih.gov/sites/entrez), enter a gene name in the search field, click the search button, and then save the page as an HTML file. Our example uses the gene vWF.# Example 4-29 shows a program for extracting some information from the results returned. The patterns it uses are very specific to results saved from Entrez Gene, but the program would be quite useful if you needed to process many such pages.

```
Example 4-29. Searching for data in an HTML file
endresults = '- - - - - - end Results - - - - - '
patterns = ('</em>]',
            '\n',
            '</a></div><div class="rprtMainSec"><div class="summary">',
def get field(contents, pattern, endpos):
    endpos = contents.rfind(pattern, 0, endpos)
    if endpos < 0:</pre>
        raise StopIteration
    startpos = contents.rfind('>', 0, endpos)
    return (endpos, contents[startpos+1:endpos])
def get next(contents, endpos):
    fields = []
    for pattern in patterns:
        endpos, field = get field(contents, pattern, endpos)
```

#vWF stands for "von Willebrand Factor," which plays a role in von Willebrand disease, the most common human hereditary coagulation abnormality. There are several forms of the disease, other genes involved, and complex hereditary patterns.

```
fields.append(field)
    fields.reverse()
    return endpos, fields
def get gene info(contents):
    lst = []
    endpos = contents.rfind(endresults, 0, len(contents))
    try:
        while(True):
            endpos, fields = get next(contents, endpos)
            lst.append(fields)
    except StopIteration:
        pass
    lst.reverse()
    return 1st
def get gene info from file(filename):
    with open(filename) as file:
        contents = file.read()
    return get_gene_info(contents)
def show_gene_info_from_file(filename):
    infolst = get_gene_info_from_file(filename)
    for info in infolst:
        print(info[0], info[1], info[2], sep='\n
                                                      ')
if __name__ == '__main__':
    show_gene_info_from_file(sys.argv[1])
                        if len(sys.argv) > 1
                        else 'EntrezGeneResults.html')
Output for the first page of the Entrez Gene results for vWF looks like this:
         Von Willebrand factor homolog
         Mus musculus
    VWF
         von Willebrand factor
         Homo sapiens
    VWF
         von Willebrand factor
         Canis lupus familiaris
         Von Willebrand factor homolog
         Rattus norvegicus
         von Willebrand factor
         Bos taurus
         von Willebrand factor
         Pan troglodytes
    VWF
         von Willebrand factor
         Macaca mulatta
```

```
vwf
    von Willebrand factor
    Danio rerio
    von Willebrand factor
    Gallus gallus
    von Willebrand factor
    Sus scrofa
    lectin
    Bombyx mori
    von Willebrand factor
    Oryctolagus cuniculus
    von Willebrand factor
    Felis catus
VWF
    von Willebrand factor
    Monodelphis domestica
    von Willebrand Factor like 2
    Ciona intestinalis
ADAMTS13
    ADAM metallopeptidase with thrombospondin type 1 motif, 13
MADE 03506
    Secreted protein, containing von Willebrand factor (vWF) type A domain
    Alteromonas macleodii 'Deep ecotype'
    putative secreted protein, containing von Willebrand factor (vWF) type A domain
    gamma proteobacterium NOR51-B
BLD 1637
    von Willebrand factor (vWF) domain containing protein
    Bifidobacterium longum DJ010A
NOR53 416
    secreted protein, containing von Willebrand factor (vWF) type A domain
    gamma proteobacterium NOR5-3
```

This code was developed in stages. The first version of the program had separate functions get symbol, get name, and get species. Once they were cleaned up and working correctly it became obvious that they each did the same thing, just with a different pattern. They were therefore replaced with a single function that had an additional parameter for the search pattern.

The original definition of get next contained repetitious lines. This definition replaces those with an iteration over a list of patterns. These changes made the whole program easily extensible. To extract more fields, we just have to add appropriate search patterns to the patterns list.

It should also be noted that because the second line of some entries showed an "Official Symbol" and "Name" but others didn't, it turned out to be easier to search backward

from the end of the file. The first step is to find the line demarcating the end of the results. Then the file contents are searched in reverse for each pattern in turn, from the beginning of the file to where the last search left off. (Note that although you might expect it to be the other way around, the arguments to rfind are interpreted just like the arguments to find, with the second less than the third.)

The Grand Unified Bioinformatics File Parser

This section explores some ways the process of reading information from text files can be generalized.

Reading the sequences in a FASTA file

Example 4-30 presents a set of functions for reading the sequences in a FASTA file. They are actually quite general, and can work for a variety of the kinds of formats typically seen in bioinformatics. The code is a lot like what we've seen in earlier examples. All that is needed to make these functions work for a specific file format is an appropriate definition of skip intro and next item.

```
Example 4-30. Reading the sequences in a FASTA file
```

```
def get items from file(filename, testfn=None):
     ""Return all the items in the file named filename; if testfn
    then include only those items for which testfn is true""
    with open(filename) as file:
        return get items(file, testfn)
def find item in file(filename, testfn=None):
    """Return the first item in the file named filename; if testfn
    then return the first item for which testfn is true"""
    with open(filename) as file:
        return find item(file, testfn)
def find item(src, testfn):
    """Return the first item in src; if testfn then return the first item for which testfn is true"""
    gen = item generator(src, testfn)
    item = next(gen)
    if not testfn:
        return item
    else:
        try:
             while not testfn(item):
                 item = next(gen)
             return item
        except StopIteration:
             return None
def get items(src, testfn=None):
     ""Return all the items in src; if testfn then include
    only those items for which testfn is true"""
    return [item for item in item generator(src)
```

```
if not testfn or testfn(item)]
def item generator(src):
    """Return a generator that produces a FASTA sequence from src each time it is called"""
    skip intro(src)
    seg = ''
    description = src.readline().split('|')
    line = src.readline()
    while line:
        while line and line[0] != '>':
             seq += line
             line = src.readline()
        yield (description, seq)
        description = line.split('|')
        line = src.readline()
def skip intro(src):
    """Skip introductory text that appears in src before the first item"""
                                     # no introduction in a FASTA file
```

The functions get items from file and find item in file simply take a filename and call get items and find item, respectively. If you already have an open file, you can pass it directly to get items or find item. All four functions take an optional filter function. If one is provided, only items for which the function returns true are included. Typically, a filter function like this would be a lambda expression. Note that find item can be called repeatedly on the same open file, returning the next item for which testfn is true, because after the first one is found the rest of the source is still available for reading.

next item is a generator version of the functions we've seen for reading FASTA entries. It reads one entry each time it is called, returning the split description line and the sequence as a pair. This function and possibly skip intro would need to be defined differently for different file formats. The other four functions stay the same.

Generalized parsing

Extracting a structured representation from a text file is known as parsing. Python, for example, parses text typed at the interpreter prompt or imported from a module in order to convert it into an executable representation according to the language's rules. Much of bioinformatics programming involves parsing files in a wide variety of formats. Despite the ways that formats differ, programs to parse them have a substantial underlying similarity, as reflected in the following template.

TEMPLATE

Grand Unified Bioinformatics File Parser

Example 4-30 defines six functions. Four of them are essentially universal, but each file format will require its own definition of next item, and if the format includes introductory text that must be skipped, skip intro will need to be redefined as well. A large proportion of bioinformatics text files can be read using this set of six functions. As a reference, an outline of what each does follows.

```
# ---- Convenience functions for starting with a filename ----
get items from file(filename, testfn=None)
find item in file(filenamee, testfn=None):
# ---- Primary functions: get all and find next ----
get items(src, testfn=None)
find item(src, testfn=None)
# ---- Format-specific functions ----
skip intro(src)
next item(src)
```

Parsing GenBank Files

Next, we'll look at an example of applying the generalized parser template to read features and sequences from GenBank flat files.* There are many ways to navigate in a browser to get a page in GenBank format from the NCBI website. For instance, if you know the GenInfo Identifier (GI), you can get to the corresponding GenBank record using the URL http://www.ncbi.nlm.nih.gov/nuccore/ followed by the GI number. Then, to download the page as a flat text file, simply click on the "Download" drop-down on the right side of the page just above the name of the sequence and select "GenBank" as the format. The file will be downloaded as sequence.gb to your browser's default download directory.

There's a great deal of information in these GenBank entries. For this example we just want to extract the accession code, GI number, feature information, and sequence. Example 4-31 shows the code needed to implement the format-specific part of the unified parser template: skip intro and next item. For a given format, the implementation of either of these two functions may require other supporting functions.

^{*} See http://www.ncbi.nlm.nih.gov/Sitemap/samplerecord.html for an example and an explanation.

[†] See http://www.ncbi.nlm.nih.gov.

```
Example 4-31. Reading sequences from a GenBank file
```

```
def get GenBank items and sequence from file(filename):
    with open(filename) as file:
        return [get ids(file), get items(file), get sequence(file)]
def get ids(src):
    line = src.readline()
    while not line.startswith('VERSION'):
        line = src.readline()
    parts = line.split()
                                                           # split at whitespace; removes \n
    assert 3 == len(parts), parts
                                                           # should be VERSION acc GI:id
    giparts = parts[2].partition(':')
    assert giparts[2], giparts
                                                           # if no colon, [1] & [2] are empty
    assert giparts[2].isdigit()
                                                           # all numbers?
    return (parts[1], giparts[2])
def get sequence(src):
     ""Return the DNA sequence found at end of src"""
    # When this is called the ORIGIN line should have just been read,
    # so we just have to read the sequence lines until the // at the end
    sea = ''
    line = src.readline()
    while not line.startswith('//'):
        seq += line[10:-1].replace(' ', '')
        line = src.readline()
    return seq
def skip intro(src):
    """Skip introductory text that appears before the first item in src"""
    line = src.readline()
    while not line.startswith('FEATURES'):
        line = src.readline()
attribute prefix = 21*' ' + '/'
def is attribute start(line):
    return line and line.startswith(attribute prefix)
def is feature start(line):
    return line and line[5] != ' '
def next item(src):
    """Return a generator that produces a FASTA sequence from src each time it is called"""
    skip intro(src)
    line = src.readline()
    while not line.startswith('ORIGIN'):
        assert is feature start(line)
                                                           # line should start a feature
        feature, line = read feature(src, line)
        # need to keep line to feed back to read_feature
        yield feature
```

```
def read feature(src, line):
    feature = line.split()
    props = {}
    line = src.readline()
    while not is feature start(line):
        key, value = line.strip()[1:].split('=')
        # remove initial / and split into [feature, value]
        if value[0] == '"':
            value = value[1:]
                                                          # remove first "; remove final " later
        fullvalue, line = read value(src, line, value)
        # need to keep line to feed back to read value
        props[key] = fullvalue
    feature.append(props)
    return feature, line
def read value(src, line, value):
    line = src.readline()
    while (not is attribute start(line) and
           not is feature start(line)):
        value += line.strip()
        line = src.readline()
    if value[-1] == '"':
        value = value[:-1]
                                                          # remove final "
    return value, line
```

The template is meant as a helpful outline, not a restrictive structure. A program written according to this template may add its own actions. In this case, the "items" to be read are the features. Before reading the features, the program extracts the accession and GI numbers. After all the features have been read, an extra step is needed to read the sequence. The top-level function returns those items in a list: a pair containing the accession and GI numbers, the list of features, and the sequence. Each feature is a list containing the type of the feature, the range of bases it covers, and a dictionary of key/ value pairs defining properties.

For the GenBank sample record saved from http://www.ncbi.nlm.nih.gov/Sitemap/sam plerecord.html, the result of executing the code would be as follows (sequence strings have been truncated to fit on a single line, and explanations have been added to the output as comments):

```
>>> data = get genbank items and sequence from file('sequences/sample.gb')
>>> pprint.pprint(data)
[('U49845.1', '1293613'),
                                                           # (accession, GI)
 [['source',
                                                          # first feature is source
    '1..5028',
                                                          # range of source within base seq
   {'chromosome': 'IX',
     db xref': 'taxon:4932',
                                                           # reference to Taxonomy DB
    'map': '9',
    'organism': 'Saccharomyces cerevisiae'}],
  ['CDS',
                                                           # coding sequence
   '<1..206',
                                                           # seq from base 1 to 206, 5' partial
   {'codon start': '3',
                                                           # translation starts at 3
     'db xref': 'GI:1293614',
                                                          # protein GI
    'product': 'TCP1-beta',
                                                          # protein produced by the CDS
```

```
'protein id': 'AAA98665.1',
                                                      # protein accession
   'translation': 'SSIYNGIS...'}],
                                                      # amino acid sequence
                                            # gene AXL2 spans nucleotides 687 through 3158
['gene', '687..3158', {'gene': 'AXL2'}],
 ['CDS',
  '687..3158',
                                                      # a complete coding sequence
  {'codon start': '1',
   'db xref': 'GI:1293615',
   'function': 'required for axial budding pattern of S.cerevisiae',
   'gene': 'AXL2',
   'note': 'plasma membrane glycoprotein',
   'product': 'Axl2p',
   'protein id': 'AAA98666.1',
   'translation': 'MTQLQISL...'}],
['gene', 'complement(3300..4037)', {'gene': 'REV7'}],
  'CDS',
  'complement(3300..4037)',
                                                      # CDS is on opposite strand
 {'codon start': '1',
   'db xref': 'GI:1293616',
   'gene': 'REV7',
   'product': 'Rev7p',
   'protein_id': 'AAA98667.1',
   'translation': 'MNRWVEKW...'}]],
                                                      # base sequence:
'gatcctccatatacaacggtatctccacctcaggtttagatctcaacaacggaaccattgcc...']
```

Translating RNA Sequences

Next, we're going to build a program to translate RNA sequences directly into strings of three-letter amino acid abbreviations. We actually got a start on this in the previous chapter, where we defined a codon table and a lookup function. They will be repeated here for convenience.

Step 1

Example 4-32 begins a series of function definitions interleaved with brief explanatory text and sample printouts.

```
RNA codon table = {
                                           Second Base
#
                                                                                       G
# U
       'UUU': 'Phe', 'UCU': 'Ser', 'UAU': 'Tyr', 'UGU': 'Cys',
                                                                                                              # UxU
      'UUC': 'Phe', 'UCC': 'Ser', 'UAC': 'Tyr', 'UGC': 'Cys', 'UUA': 'Leu', 'UCA': 'Ser', 'UAA': '---', 'UGA': '---', 'UUG': 'Leu', 'UCG': 'Ser', 'UAG': '---', 'UGG': 'Trp',
                                                                                                              # UxC
                                                                                                              # UxA
                                                                                                              # UxG
       'CUU': 'Leu', 'CCU': 'Pro', 'CAU': 'His', 'CGU': 'Arg',
                                                                                                              # CxU
       'CUC': 'Leu', 'CCC': 'Pro', 'CAC': 'His', 'CGC': 'Arg', 'CUA': 'Leu', 'CCA': 'Pro', 'CAA': 'Gln', 'CGA': 'Arg', 'CUG': 'Leu', 'CCG': 'Pro', 'CAG': 'Gln', 'CGG': 'Arg',
                                                                                                              # CxC
                                                                                                              # CxA
                                                                                                              # CxG
# A
       'AUU': 'Ile', 'ACU': 'Thr', 'AAU': 'Asn', 'AGU': 'Ser',
                                                                                                              # AxU
       'AUC': 'Ile', 'ACC': 'Thr', 'AAC': 'Asn', 'AGC': 'Ser', 'AUA': 'Ile', 'ACA': 'Thr', 'AAA': 'Lys', 'AGA': 'Arg', 'AUG': 'Met', 'ACG': 'Thr', 'AAG': 'Lys', 'AGG': 'Arg',
                                                                                                              # AxC
                                                                                                              # AxA
                                                                                                              # AxG
       'GUU': 'Val', 'GCU': 'Ala', 'GAU': 'Asp', 'GGU': 'Gly',
                                                                                                              # GxU
       'GUC': 'Val', 'GCC': 'Ala', 'GAC': 'Asp', 'GGC': 'Gly', 'GUA': 'Val', 'GCA': 'Ala', 'GAA': 'Glu', 'GGA': 'Gly', 'GUG': 'Val', 'GCG': 'Ala', 'GAG': 'Glu', 'GGG': 'Gly'
                                                                                                              # GxC
                                                                                                              # GxA
                                                                                                              # GxG
}
def translate RNA codon(codon):
      return RNA codon table[codon]
```

The next step is to write a function that translates an RNA base string into a string of the corresponding three-letter amino acid abbreviations. The optional step argument to range is useful for this. Testing with assertions while this code was being developed revealed the need to ignore the last base or two of sequences whose length is not a multiple of 3, something not considered when the code was first written. The expression len(seq)%3 gives the remainder when the length of the sequence is divided by 3—we have to subtract that from len(seq) so we don't try to process an excess base or two at the end of the sequence. The new example is shown in Example 4-33.

Example 4-33. Translating RNA sequences, step 2

```
def translate(seq):
    """Return the animo acid sequence corresponding to the RNA sequence seq"""
    translation = ''
    for n in range(0, len(seq) - (len(seq) % 3), 3):
                                                                     # every third base
        translation += translate RNA codon(seq[n:n+3])
    return translation
```

Next, we take care of frame shifts and add printing functions with the functions shown in Example 4-34.

```
Example 4-34. Translating RNA sequences, step 3
def translate in frame(seg, framenum):
    """Return the translation of seq in framenum 1, 2, or 3"""
    return translate(seq[framenum-1:])
def print translation in frame(seq, framenum, prefix):
    """Print the translation of seq in framenum preceded by prefix"""
        print(prefix,
          framenum,
           ' * framenum,
          translate in frame(seq, framenum),
def print translations(seq, prefix=''):
    """Print the translations of seq in all three reading frames, each preceded by prefix"""
    print('\n' ,' ' * (len(prefix) + 2), seq, sep='')
    for framenum in range(1,4):
        print translation in frame(seq, framenum, prefix)
>>> print translations('ATGCGTGAGGCTCTCAA')
 ATGCGTGAGGCTCTCAA
1 MetArgGluAlaLeu
2 CysValArgLeuSer
3 Ala---GlySerGln
>>> print translations('ATGATATGGAGGAGGTAGCCGCGCCCATGCGCGCTATATTTTGGTAT')
  ATGATATGGAGGAGGTAGCCGCGCGCCATGCGCGCTATATTTTGGTAT
1 MetIleTrpArgArg---ProArgAlaMetArgAlaIlePheTrpTyr
2 ---TyrGlyGlyGlySerArgAlaProCysAlaLeuTyrPheGly
  AspMetGluGluValAlaAlaArgHisAlaArgTyrIleLeuVal
```

Step 4

Now we are ready to find the open reading frames. (We make the simplifying assumption that we're using the standard genetic code.) The second and third functions here are essentially the same as in the previous step, except that they call translate with open reading frames instead of translate in frame. Example 4-35 shows the new definitions.

Example 4-35. Translating RNA sequences, step 4

```
def translate with open reading frames(seq, framenum):
    """Return the translation of seq in framenum (1, 2, or 3), with ---'s when not within an
    open reading frame; assume the read is not in an open frame when at the beginning of seq"""
    open = False
    translation = ""
    seqlength = len(seq) - (framenum - 1)
    for n in range(frame-1, seqlength - (seqlength % 3), 3):
        codon = translate RNA codon(seq[n:n+3])
```

```
open = (open or codon == "Met") and not (codon == "---")
       translation += codon if open else "---"
   return translation
def print translation with open reading frame(seq, framenum, prefix):
   print(prefix,
         framenum,
         ' ' * framenum,
         translate with open reading frames(seq, framenum),
         sep='')
def print translations with open reading frames(seq, prefix=''):
   print('\n', ' ' * (len(prefix) + 2), seq, sep='')
   for frame in range(1,4):
       print_translation_with_open_reading_frame(seq, frame, prefix)
>>> print translations('ATGCGTGAGGCTCTCAA')
 ATGCGTGAGGCTCTCAA
1 MetArgGluAlaLeu
>>> print translations('ATGATATGGAGGAGGTAGCCGCGCCCATGCGCGCTATATTTTGGTAT')
 ATGATATGGAGGAGGTAGCCGCGCGCCATGCGCGCTATATTTTGGTAT
1 MetIleTrpArgArg------MetArgAlaIlePheTrpTyr
2 ------
3 ---MetGluGluValAlaAlaArgHisAlaArgTyrIleLeuVal
```

Finally, we print the sequence both forward and backward. Getting the reverse of a sequence is easy, even though there's no function for it: seq[::-1]. Remember that trick, as you will need it any time you want to reverse a string. Working with biological sequence data, that will be quite often! Example 4-36 shows the final piece of the code.

```
Example 4-36. Translating RNA sequences, step 5
def print translations in frames in both directions(seq):
   print translations(seq, 'FRF')
   print translations(seq[::-1], 'RRF')
def print translations with open reading frames in both directions(seq):
   print translations with open reading frames(seq, 'FRF')
   print translations with open reading frames(seq[::-1], 'RRF')
>>> print translations('ATGCGTGAGGCTCTCAA')
    ATGCGTGAGGCTCTCAA
FRF1 MetArgGluAlaLeu
FRF2 -----
FRF3 -----
    AACTCTCGGAGTGCGTA
RRF1 -----
RRF3
     -----
```

```
>>> print translations('ATGATATGGAGGAGGTAGCCGCGCCCATGCGCGCTATATTTTGGTAT')
    ATGATATGGAGGAGGTAGCCGCGCGCCATGCGCGCTATATTTTGGTAT
FRF1 MetIleTrpArgArg------MetArgAlaIlePheTrpTyr
     ---MetGluGluValAlaAlaArgHisAlaArgTyrIleLeuVal
    TATGGTTTTATATCGCGCGTACCGCGCGCGCGATGGAGGAGGTATAGTA
RRF2 MetValLeuTyrArgAlaTyrArgAlaProMetGluGluVal---
RRF3
```

Constructing a Table from a Text File

Our next project will be to construct a table from a text file. We'll use the file located at http://rebase.neb.com/rebase/link bionet, which contains a list of restriction enzymes and their cut sites. The file has a simple format.[‡] Each line of data has the form:

```
EnzymeName (Prototype)
                             ... spaces ...
```

A "prototype" in this data set is the first enzyme to be discovered with the specified cut site. Lines that represent prototypes do not have anything in the "Prototype" column.

Several lines of information appear at the beginning of the file, before the actual data. To ensure that our program ignores these lines we'll use one of our usual skip introtype functions, here called get first line. A look at the file shows that the first line of data is the first one that begins with an A. This is certainly not an acceptable approach for "production" software, since the organization of the file might change in the future, but it's good enough for this example. The end of the file may have some blank lines, and we'll need to ignore those too.

To represent the data in this file we'll construct a dictionary whose keys are enzyme names and whose values are the cut sites. We'll make this simple and ignore the information about prototypes. Because there are so many details we are going to take things a step at a time. This is how you should work on your programs too.

Step 1

The general outline of the program will be:

- 1. Initialize the enzyme table.
- 2. Skip introductory lines, returning the first real line.
- 3. While line is not empty:
 - a. Parse line.
 - b. Store entry in the enzyme table.
 - c. Read another line.

[#] The site contains files in other formats with more information. We'll use one of those later in the book.

Turning those steps into function names and adding as few details as we can get away with, we write some essentially empty functions (see Example 4-37). For example, get first line just returns an empty string; in a sense it's done its job, which is to return a line.

```
Example 4-37. Simple Rebase reader, step 1
def load enzyme table():
    return load enzyme data into table({})
    # start with empty dictionary
def load enzyme data into table(table):
    line = get first line()
    while not end of data(line):
        parse(line)
        store entry(table)
        line = get next line()
    return table
def get first line():
    return ''
                                                    # stop immediately
def get next line():
    return ' '
                                                    # so it stops after getting the first line
def end of data(line):
    return True
def parse(line):
    return line
def store entry(table):
    pass
# testing:
def test():
    table = load enzyme table()
    assert len(table) == 0
    print('All tests passed.')
test()
```

Step 2

We can fill in the details of some of these functions immediately. Do not be disturbed that some of the definitions remain trivial even after all the changes we'll make. We might want to modify this program to use with a different file format, and there's no guarantee that get next line, for instance, will always be as simple as it is here. Using all these function names makes it very clear what the code is doing without having to comment it.

In the following steps, changes from and additions to the previous step are highlighted. They include:

- Binding names to the result of parse (using tuple unpacking)
- Passing the key and value obtained from parse to store entry
- Returning a result from first line to use in testing
- Implementing end of data
- Splitting the line into fields and returning the first and last using tuple packing
- Implementing store entry

Many of the functions continue to have "pretend" implementations in step 2, which is shown in Example 4-38.

```
Example 4-38. Simple Rebase reader, step 2
def load enzyme table():
    return load enzyme data into table({})
    # start with empty dictionary
def load enzyme data into table(table)
    line = get first line()
    while not end of data(line):
        key, value = parse(line)
        store entry(table, key, value)
        line = get next line()
    return table
def get first line():
    return 'enzymeA (protoA)
                                            CCCGGG'
    # return a typical line
def get next line():
    return '-'
                                        # so it stops after getting the first line
def end of data(line):
    return len(line) < 2
                                        # 0 means end of file, 1 would be a blank line
def parse(line):
    fields = line.split()
                                        # with no argument, split splits at whitespace
                                        # tuple packing (omitting optional parens)
    return fields[0], fields[-1]
                                        # avoiding having to determine whether there are 2 or 3
def store entry(table, key, value):
    table[key] = value
def test():
    table = load enzyme table()
    assert len(table) == 1
    result = parse('enzymeA (protoA)
                                                      CCCGGG')
```

```
assert result == ('enzymeA', 'CCCGGG'), result
    print('All tests passed.')
test()
```

In the next step, we actually read from the file. It's silly to try to wrestle with a large datafile while you are writing the functions to handle it. Instead, extract a small bit of the actual file and put it in a "test file" that you can use until the program seems to work. We'll construct a file called rebase_test_data01.txt that contains exactly the following text:

```
some introductory text
more introductory text
AnEnzyme (APrototype)
                                  cutsite1
                                  cutsite2
APrototype
```

Changes in this step include making some of the definitions more realistic:

- Adding a filename parameter to load enzyme table
- Embedding most of that function in a loop
- Adding a call to print for debugging purposes
- Passing the open file object to return_first_line and get_next_line
- Implementing get first line and get next line

Example 4-39 illustrates the third step.

```
Example 4-39. Simple Rebase reader, step 3
def load enzyme table(data filename):
    with open(data filename) as datafile:
        return load enzyme data into table(datafile, {})
def load enzyme data into table(datafile, table)
    line = get first line(datafile)
    while not end of data(line):
        print(line, end='')
        key, value = parse(line)
        store entry(table, key, value)
        line = get next line(datafile)
    return table
def get first line(fil):
     line = fil.readline()
     while line and not line[0] == 'A':
         line = fil.readline()
    return line
def get next line(fil):
    return fil.readline()
```

```
def end of data(line):
    return len(line) < 2
def parse(line):
    fields = line.split()
    return fields[0], fields[-1]
def store entry(table, key, value):
    table[key] = value
def test():
    print()
    datafilename = 'rebase test data01.txt'
    table = load enzyme table(datafilename)
    assert len(table) == 2, table
    result = parse('enzymeA (protoA)
                                                  CCCGGG')
    assert result == ('enzymeA', 'CCCGGG'), result
    print('All tests passed.')
test()
```

Finally, we clean up some of the code (not shown here), use the real file, and test some results. Example 4-40 shows step 4.

```
Example 4-40. Simple Rebase reader, step 4
```

This step uses the definitions of the previous step unchanged, except # that the call to print in load_enzyme_data_into_table could be removed

```
def test():
    print()
    datafilename = 'link bionet.txt'
    table = load enzyme Table(datafilename)
    # check first entry from file:
    assert table['AaaI'] == 'C^GGCCG'
    # check an ordinary entry with a prototype:
    assert table['AbaI'] == 'T^GATCA', table
    # check an ordinary entry that is a prototype:
    assert table['BclI'] == 'T^GATCA', table
    # check last entry from file:
    assert table['Zsp2I'] == 'ATGCA^T'
    assert len(table) == 3559, len(table)
    print()
    print('All tests passed.')
```

Step 5

If we wanted to, we could also add a function for printing the table in a simpler format to make it easier to read in the future and a corresponding function for doing that reading. We can print each entry of the table on a separate line with the name of the

enzyme separated by a tab from the sequence it recognizes. Example 4-41 shows these two simple functions.

```
Example 4-41. Simple Rebase reader, step 5
def write table to filename(table, data filename):
    """Write table in a simple format to a file named data_filename"""
    with open(data filename, 'w') as file:
        write table entries(table, files)
def write table entries(table, datafile):
    for enzyme in sorted(table.keys()):
        print(enzyme, table[enzyme], sep='
                                                     ', file=datafile)
def read table from filename(data filename):
    ""Return a table read from the file named data filename
    that was previously written by write_table_to_filename"""
    with open(data filename) as file:
        return read table entries(file, {})
def read table entries(datafile):
    for line in datafile:
        fields = line.split()
        table[fields[0]] = fields[1]
    return table
```

Tips, Traps, and Tracebacks

Tips

- Keep functions small.
- Prefer comprehensions to loops or iterations that collect values.
- Look for opportunities to use conditional expressions—(val1 if val2 else val3) —instead of if/else statements when the two clauses contain just an expression or assign the same name.
- The mechanisms discussed in this chapter are the core of Python programs. You should review the chapter from time to time to better understand parts you didn't completely follow the first time through.
- Use the templates as a reference—they capture a large portion of the ways control statements are used.
- In general, if a function is to return True if a certain condition holds and False otherwise, spelling out the following is technically "silly":

```
if condition:
    return True
else:
   return False
```

Instead, just write return condition. For example, instead of:

```
if len(seq1) > len(seq2):
        return True
    else:
        return False
write:
    return len(seq1) > len(seq2)
```

The result of the comparison is True or False.

• When it is last in the function, a statement such as the following (without an else clause):

```
if boolean-condition:
        return expression
can be more concisely written as:
    return boolean-condition and expression
```

This expression will return False if condition is false and the value of expression if it is true. The primary reason you would need to use the conditional statement instead of expression is if you specifically want the function to return None if condition is false, rather than False.

- While assertions are valuable for testing code and for checking conditions at certain points in function definitions while the definitions are still under development, do not use assertions as a coding technique in place of conditionals. For example, if you want to check the validity of some input, do that in a conditional statement and take appropriate action if the condition is violated, such as raising an error. Do not simply assert that condition.§
- The first Python code in a file should be a docstring. If enclosed in triple single or double quotes, the docstring can be as many lines long as you want. If your file is imported (rather than executed), the module that will be created for it will store that docstring, and Python's help facility will be able to display it. You'll probably want to comment the file to explain what it is, even if only for your own future reference, so you might as well get in the habit of using docstrings rather than comments.
- You should make a choice about how you will use single single, single double, triple single, and triple double quotes, then follow that convention in a consistent way. The reason for using each of the four kinds of quotes in a different way is that it makes it easier to search for appearances in your code of a string that serves a particular purpose, such as a docstring.

[§] Other than coding style, the issue here is that assertions are meant to be executed only during development, not in a "production" version of a program. When running Python from the command line, the -0 option can be added to optimize a few aspects of the execution, one of which is to ignore assertions.

The following choices were made for the code in this book and the downloadable code files:

Single single quotes

Used for short strings

Single double quotes

Used for short strings with internal single quotes

Triple double quotes

Used for docstrings

Triple single quotes

Used for long value strings

The difference between single single quotes and single double quotes is not so important, but it's better to use one most of the time. Don't forget that you can include a single quote inside a single-quoted string (or a double quote inside a double-quoted string) simply by putting a backslash before it.

- If a compilation error points to a line that appears to be OK, the problem is probably on the previous line. Check that all "structural" lines of compound statements def, if, else, elif, while, for, try, except, and finally—end with a colon.
- When your IDE indents a line to an unexpected level, don't fight it or ignore it it's giving you very useful information about a problem in your code. Make sure that each colon in a compound statement is at the end of its "logical line" (i.e., that the beginning and any continuation lines are treated as one). The only keywords that are followed by a colon (and therefore nothing else on that line) are else, try, and finally.
- Here's a useful debugging function. It is better than just using assert for testing code because an assertion failure stops the program, whereas failures identified by calls to this function do not:

```
def expect equal(expected, result):
    """Return expected == result, printing an error message if not true;
    use by itself or with assert in a statement such as:
        assert expect equal(3438,
                               count hypothetical proteins(gbk filename)"""
    if expected == result:
        return True
    print('Expected', expected, 'but got', result)
```

Traps

 Many functions that collect values, whether using a loop or an iteration, must avoid adding to the collection on the first pass through the loop or iteration. When that is the case, the loop or iteration must be followed by a statement that adds the last item to the collection.

- When processing data read from outside the program, do not trust that all of it is in the expected form. There may be missing or aberrant lines or entries; for example, an enzyme cut site that should be? to mean "unknown" may be the digit 7 instead.
- It is rarely a good idea to process large files by calling readlines. Rather, process one line at a time, by calling readline in a loop, by using a comprehension over the lines of the file, or by using a for statement. This avoids creating enormous lists that occupy large amounts of memory.

Tracebacks

Following are some representative error messages:

AttributeError: 'range' object has no attribute 'index'

A call has been made to a method not supported by the type.

F0FFrror

The programmer (or user) typed Ctrl-D (Ctrl-Z on Windows).

IndentationError: unexpected indent

This can happen (well) after a try statement with no except or finally clause.

IOError: [Errno 2] No such file or directory: 'aa2.fasta'

There are a number of IOError variations, each with a different Errno and message format.

KeyboardInterrupt

The user typed Ctrl-C twice.

KeyError

An attempt has been made to reference a dictionary element by a key not present in the dictionary.

TypeError: unorderable types int() < str()</pre>

An attempt has been made to compare two values of different types.

TypeError: 'NoneType' object is not iterable

The code contains a for statement that is iterating over something that has no value. This is *not* the same as iterating over an empty collection, which, although it does nothing, does not cause an error. Errors that mention NoneType are almost always caused by a function meant to return a value that does not include a return statement.

object is not iterable: 'builtin_function_or_method'

A function name was used in place of a function call in the in part of a comprehension or for statement; i.e., you forgot the parentheses. This is a common mistake when calling dict.keys(), dict.values(), and dict.items().



Classes

Simple collections are appropriate for organizing relatively simple and transient information. Elaborately nested collections, though, are not. Consider the output parsed from the very small GenBank entry shown in Example 4-31. Even though it ignores much of the file's information, the program produces an elaborate nested structure. Working with such a structure creates the following serious problems:

Complexity

Using the representation in a program requires understanding its intricacies.

Awkward navigation

Accessing specific parts of the representation requires tricky combinations of index expressions and function calls.

Exposure to change

Changes to the representation require widespread changes in every program that uses it

The combination of these factors causes mental strain, slows programming, makes debugging more difficult, and increases mistakes. It is too easy to write [1] instead of [0], or [1] instead of [0][1]. Later, such index expressions have no apparent meaning, so they are difficult to understand and correct. If changes must be made it is difficult to make them consistently and find every place that must be edited. While collections are powerful tools, they are best reserved for actions that operate on the entire collection, rather than being used with access expressions or functions that pick out specific elements.

These problems arise because of the entanglement of code that does need to know about representation details and code that does not. For instance, we can define a function called <code>get_organism</code> that extracts the name of the organism from the complex collection. That function needs to know the details of the representation. However, other functions can just call <code>get_organism</code> when they need that piece of the collection. With this approach, if the structure of the collection is changed in a way that affects

the way an organism's name is accessed, only the get organism function will be affected, not any of the code that calls it.

Functions like get organism hide details from code that uses a representation. Only a small number of functions need to work with the representation directly; everything else calls those functions. This strategy defines a functional interface to the complex structure. Changes to the representation cause changes only to the small number of functions in the interface. This enforces the principle of information hiding that dictates the separation of implementation and use.

We have already seen two kinds of information hiding:

- Accessing built-in Python types through functions and methods without knowing anything about how they are represented
- Dividing code into small chunks, with higher-level functions that organize the computation separated from lower-level ones dealing with the details of file formats and collection structures

The comments in the preceding paragraphs demonstrate the need for another kind:

 Packaging together a representation along with the functions that manipulate it so that code can call those functions rather than accessing the data

If we had the third kind of information hiding, the result would be much like the first. What we need, then, is a way to define new types (classes) the way we can define new functions. Python does indeed give us such a mechanism.

Defining Classes

Python's collection types work as follows:

- 1. The type is called like a function to create new instances.
- 2. The call to the type can include arguments used to initialize the new instance.
- 3. Methods are invoked on instances of types instead of accessing their internal representations.
- 4. Methods defined for different types can have the same name: when a method is called through a value, Python finds the appropriate function by looking in the definition of the value's type.
- 5. The way an instance is printed is based on its type.

None of this is special to collection types. These features are intrinsic to Python's type mechanisms. They are what make Python an object-oriented programming language.

Python's class definition statement defines a new class (type), just the way a definition statement defines a new function. The keyword that introduces the statement is class. Like function definition statements, class definition statements are one-clause compound statements containing other statements. The other statements are nearly always function definitions, but any statement is allowed. Each def statement inside a class defines a method for the instances of that class.

New instances are created by calling the name of the class as a function, just as with built-in types. Each instance stores information in named *fields*. Together, fields and methods are called attributes. They are accessed with the usual dot notation, such that this expression:

```
obj.name()
```

would call the name method defined by obj's type, and this expression:

```
obj.species
```

would access the species attribute of obj. That attribute might be a field or it might be a method. Using str as an example:

```
>>> 'AAU'.isupper
                                             # ACCESS the isupper ATTRIBUTE
<built-in method isupper of str object at 0x1c523e0>
>>> 'AAU'.isupper()
                                             # CALL the isupper METHOD
True
```

What classes should your programs define? They can define pretty much whatever you can name. Classes add nouns to the language you are building with your code, the way functions add verbs. The definition of a class defines the functional interface to the data its instances represent. Classes are the mechanism that supports information hiding.

Collections serve two very different purposes: grouping and structuring. A group has no intrinsic meaning—it just collects values together. A complex collection structure is different. The data returned by the GenBank parser of Example 4-31 (see "Parsing GenBank Files" on page 148) is a list, but it's not the same kind of list as one containing primitive values. It has many levels of structure, with many elements playing specific roles in what the collection represents. Classes should be used in place of all but the simplest structuring collections.

STATEMENT

Class Definition

A class definition begins with the keyword class and a name, followed by a colon and one or more statements indented relative to the first line.

```
class name:
   def method1(args...):
    def method2(args...):
```



You should begin each syllable of your class names with an uppercase letter. Omit the underscores that would otherwise separate syllables in the usual Python naming convention.

The statement in Example 5-1 defines a new Python type called GenBankEntry. Using pass for the class body is just like using pass for the body of a function you know you want to define but haven't yet determined how to implement.

Example 5-1. Defining a class

```
class GenBankEntry:
   pass
>>> GenBankEntry()
<__main__.GenBankEntry object at 0x1cafdb0>
>>> GenBankEntry()
< main .GenBankEntry object at 0x1caff30>
```

The interpreter's output when a new instance of GenBankEntry is created shows the name of its module (main , indicating the interpreter), the name of the class, and a hexadecimal memory address. The memory addresses don't mean anything, but they at least show that we have created two different instances of the class.

Values may be stored in fields of an instance using dot notation. We'll do this initially for demonstration purposes only—in general, we'll want to use a functional interface:

```
>>> gb = GenBankEntry()
>>> gb.accession = 'U49845.1'
>>> gb.gi = '1293613'
>>> gb.accession
'U49845.1'
```

Instance Attributes

To demonstrate important features of classes, instances, and methods, we'll build a class to represent the information parsed from a GenBank entry as shown in Example 4-31. The code will grow rather long as more methods are added, so only the new methods are shown for each example.

Access methods

The first kind of methods we will consider are access methods, which return values based on the fields of an instance. The simplest just return the value of a field. We want programs using our class to call methods with expressions, such as gb.get gi(), instead of directly accessing the fields with expressions such as gb.gi. To support that, we need to define the method get gi() inside the class. It will return the value of the gi field for the instance on which it is called.

How does a method determine which instance's fields to access? Python automatically adds the instance through which a method is called to the beginning of the call's argument list. Therefore, the instance a method uses is the value of its first parameter. By convention, the name of that parameter is self:

```
class GenBankEntry:
    def get gi(self):
        return self.gi
    def get accession(self):
        return self.accession
>>> gb.get accession()
'U49845.1'
```

Suppose we decide to change the way we store the accession and GI values. We might want to move them into a dictionary called info, anticipating the addition of more keys and values. It would then be necessary to redefine the methods based on this new representation:

```
class GenBankEntry:
    info = \{\}
    def get gi(self):
        return self.info['gi']
    def get accession(self):
        return self.info['accession']
```

Any code that uses the access functions would be completely unaffected by this change. This is the essence of information hiding—separating the implementation of a representation from its use:

```
>>> gb = GenBankEntrv()
>>> gb.info = {'accession': 'U49845.1', 'gi': '1293613'}
>>> gb.get_accession()
'U49845.1'
```

Now suppose we extract the source information from the data returned by the GenBank file parser and store it in a field called source. The source is the first feature in the data's list of features, which itself is the second element of the list returned. We'll store the source in one field and a list of the rest of the features in another:

```
>>> data = get GenBank items and sequence from file(genbank file name)
                  # defined in Example 4-31
>>> gb.source = data[1][0]
>>> gb.features = data[1][1:]
>>> pprint.pprint(gb.source)
['source',
 '1..5028',
 {'chromosome': 'IX',
  'db_xref': 'taxon:́4932',
```

```
'map': '9',
'organism': 'Saccharomyces cerevisiae'}]
```

We don't want outside code digging around in the source's qualifier dictionary, or even knowing where it is and that it is a dictionary. So, we'll define access methods to get values from the qualifier dictionary. We will *not* define a method that returns the value of the source field itself, since that would expose the representation of that part of the information:

```
class GenBankEntry:
    def organism(self):
        return self.source[2].get('organism', None)
    def chromosome(self):
        return self.source[2].get('chromosome', None)
```

Not all access methods return data that is directly stored in the instance. Some perform numerical computations, string manipulations, or other operations using the stored data:

```
class GenBankEntry:
    def gc content(self):
        return round((100 * ((self.sequence.count('g') +
                               self.sequence.count('c'))) /
                      len(self.sequence)),
                     2)
>>> gb.gc content()
37.97
```

Other methods do some kind of filtering or searching to find the desired information. It isn't even necessary for the data a method filters or searches to be the value of one of the instance's fields; for example, a method might search a collection obtained by calling an access method of a different instance. Here is the definition of one that filters and extracts the instance's feature list to produce a list of genes and the range of their coding region(s):

```
def genes(self):
        return [(feature[1], feature[2]['gene'])
                for feature in self.features
                if feature[0] == 'gene']
>>> gb.genes()
[('687..3158', 'AXL2'), ('complement(3300..4037)', 'REV7')]
```

TEMPLATE

Access Methods

Access methods return values based on the fields of an instance. They can do much more than just return the value of a field. Part of information hiding, in fact, is that

programmers using access methods don't have to know which ones refer to similarly named fields. Access methods generally perform one of the following kinds of computations:

- Get the value of a field
- Look up a value in a dictionary that is the value of a field
- Compute a value from the value of one or more fields
- Filter the collection that is the value of a field
- Search for a targeted value in a field that has a collection value or in some other collection outside the instance

Predicate methods

A predicate is a Boolean-valued method or function. A predicate method is essentially an access method that also does a comparison. Classes should define methods for predicates that are likely to be important, difficult to compute, or frequently used. It is better to define predicate methods for other code to use than it is to leave the other code to interpret the meaning of values returned from access methods.

To highlight that a method is a predicate, its name should usually begin with is (we've already seen quite a few predicates defined by the str class, such as isupper). Occasionally, however, the grammar of the word(s) used for the method name calls for the prefix has. If the rest of the method name has multiple syllables you would normally include an underscore after the prefix, beginning the name with is or has. A test for whether a GenBankEntry contains a base sequence or an amino acid sequence would probably be the most useful predicate for our class:

class GenBankEntry:

```
def is base sequence(self):
    return set(self.get sequence().lower()) <= {'a', 'c', 't', 'g'}</pre>
```

One predicate that it is often important to provide is one that determines whether an instance is before another. This would be used to implement some kind of ordering among the instances of the class. The method 1t is called by the built-in function sorted and the method list.sort. Any call to one of these can include a key argument to specify an ad hoc sorting order. If no key argument is supplied in a call, 1t is used.



Method names like 1t that begin and end with two underscores are called special methods. They play predefined roles in Python's implementation. You are not allowed to give other methods names that begin and end with two underscores.

Many classes should implement 1t, even if all it does is compare the name or identifier of its two arguments. For a class that does not implement 1t, it is an error to try to sort a sequence of its instances. The __lt__ method must have two parameters: self and another for the other object to which self will be compared. If we want the method to be able to compare only objects of the same type, we can begin its definition by testing whether they are of the same type and raising an exception if not. Here's a definition of __lt__ that compares two instances of GenBankEntry using their GenInfo IDs:

Though limited to comparing two instances of the same type, that definition is general enough to capture it as a template.

TEMPLATE

Supporting Sorting

The method _lt_ is called by sorted and list.sort when no key argument is provided. A simple definition would check that its two arguments have the same type; more sophisticated definitions could handle multiple types.

```
class ClassName:
```

Initialization methods

One special method, __init__, is critical to the way classes work. When a class is called to create a new instance, the arguments included in the call are passed automatically to the __init__ method defined in the class. This method can initialize the instance's fields and ensure that the new instance is ready for use. This is how fields are normally assigned their initial values, not by using dot notation with the field name outside of the class.

The __init__ method plays a fundamental role in the interface defined by the class. Although other methods could add other fields, the norm is to assign all fields in a class's __init__ method, even if the value is only None. To help anyone reading the class, including its author, the __init__ method should go first in the body of the class's definition.

Following is the definition of a possible init method for the GenBankEntry class. It is designed to take just one argument other than self: the full complex of data read from the GenBank entry parser. Note that init methods embody the choices the class's author has made about field names and what goes in which fields. Because code outside the class would use access methods, not field names, no one using the class needs to know about anything inside the init method:

class GenBankEntry:

```
def init (self, data):
    self.accession, self.version = data[0][0].split('.')
    self.gid = data[0][1]
    self.sequence = data[2]
    # assuming first feature is source & treating specially
    assert 'source' == data[1][0][0]
    self.features = data[1][1:]
    self.source = data[1][0]
```

Now we no longer need to assign fields outside of the class. Instead, we just call the class with the data obtained from the parser:

```
>>> gb = GenBankEntry(data)
```

This creates a new instance, calls GenBankEntry.__init__ (passing it the new instance as the first argument and data as the second), and assigns gb to the result. In this one case, there is no instance in front of a dot in the method call: Python automatically and invisibly calls the init method as part of the instance creation process. By the time it calls init, the new instance has been created, so it can be passed as the self argument to the method. Note that the init method does not return a value. After the method is called, the new instance is returned as the value of the call to the class.

String methods

Two other important special methods are str and repr . These define how Python prints instances of a class. Each returns a string. What that string contains is up to the class implementer.

The two methods are slightly different, as shown in Table 5-1. If a class defines _repr__ but not __str__, __repr__ will be called in place of __str__. However, if a class defines str but not repr , str will not be called—the default definition of repr will be used instead.

Table 5-1. __str__ versus __repr__

	str	repr
Called by	str(<i>obj</i>)	repr(obj)
Returns	A human-readable string	A string that could be typed at the interactive prompt or included in a Python file to create an equivalent object
Where "called by" functions are called	print	Interpreter output

The "called by" functions call the methods. The interpreter and the print function call the "called by" functions to get a string, then print it without the quotes. With neither str nor repr defined yet, the four possible outputs would be as follows:

```
>>> gb
<_main_.GenBankEntry object at 0x1c69db0>  # default repr, no quotes
>>> repr(gb)
'<_main_.GenBankEntry object at 0x1c69db0>'  # default repr, quotes
>>> print(gb)
<_main_.GenBankEntry object at 0x1c69db0>  # default repr, no quotes
>>> str(gb)
'<_main_.GenBankEntry object at 0x1c69db0>'  # default repr, quotes

Here is a simple definition of GenBankEntry.__str__:
```

class GenBankEntry:

```
def __str__(self):
    return 'GenBankEntry-' + self.get_gid()
```

With that definition in place, here's what happens with the same four expressions:

```
>>> gb
<__main__.GenBankEntry object at 0x1c69db0>  # default repr, no quotes
>>> repr(gb)
'<__main__.GenBankEntry object at 0x1c69db0>'  # default repr, quotes
>>> print(gb)
GenBankEntry-1293613  #__str__, no quotes
>>> str(gb)
'GenBankEntry-1293613'  #__str__, quotes
```

There are really two kinds of _repr_ implementations. For classes whose instances can be represented succinctly, such as range, the goal is to have the interpreter output be the same as the input:

```
>>> range(1,5) range(1,5)
```

For other classes, the goal is more pragmatic. We could define the first kind of _repr__ method for GenBankEntry, as it contains all the information needed to reconstruct the complex collection structure that was passed to its initialization method. That wouldn't be a good idea, however: any time a GenBankEntry was typed to the interpreter the resulting printout would be very long. That doesn't necessarily mean we must put up with the uninformative default, though. We can define __repr__ in a way that is convenient for interaction with the interpreter, debugging, and so on.

If we change the name of the __str__ method to __repr__, the same method will be called in all four of our situations, since __str__ defaults to __repr__ if not defined:

```
{\tt class} \ {\tt GenBankEntry:}
```

This is what happens with the four expressions now:

```
>>> gb
'GenBankEntry-1293613'
                                                          # __repr__, no quotes
>>> repr(gb)
'GenBankEntry-1293613'
                                                          # __repr__, quotes
>>> print(gb)
GenBankEntry-1293613
                                                          # __repr__, no quotes
>>> str(gb)
'GenBankEntry-1293613'
                                                          # __repr__, quotes
```

Finally, we also have the option of defining both methods. It is often useful to define repr to give more information than str . Usually you'll want to keep the result of _str_ fairly compact, since it is called by print. Let's define the two methods as follows, then run our expressions again:

```
def __str__(self):
   return "GenBankEntry-" + self.get gid()
def repr (self):
   return "<GenBankEntry {} {} '{}'>".format(self.get gid(),
                                             self.get accession(),
                                             self.organism()
```

```
>>> gb
<GenBankEntry 1293613 U49845.1 'Saccharomyces cerevisiae'>
>>> repr(gb)
"<GenBankEntry 1293613 U49845.1 'Saccharomyces cerevisiae'>"
>>> print(gb)
GenBankEntry-1293613
>>> str(gb)
```



'GenBankEntry-1293613'

class GenBankEntry:

These definitions of str and repr do not directly access fields of the instances. Information hiding is just as useful inside a class as outside. Generally, only a small number of methods need direct access to fields; the others can use those. That allows decisions about field names and contents to be changed without requiring extensive editing of the other methods of the class.

Modification methods

In many cases, once an instance has been created many of a class's fields will not be changed. As we've defined it so far, GenBankEntry is completely static—none of the values of an instance's fields will ever change. To make it possible for a program to change the value of a field without violating the information hiding principle, the class must define a modification method for that field.

Stretching our example just a bit, suppose we are writing a program that manages a number of GenBankEntry instances for its user, and we want to add a field to contain

notes the user makes. Since the user may make multiple notations, we'll store them in a list. We need to do three things to implement this:

- 1. Assign the field to a new list in init .
- 2. Define an access method to retrieve the notes.
- 3. Define a modification method to add a note.

As with access methods, there are different kinds of modification methods that you can use, depending on the type of the field's value. The method may just replace the primitive value of a field, or it may add elements to or remove elements from a field's collection value, navigate into a more complex field value, change multiple fields, and so on. For this example, let's assume that each note is a string, and all that happens when a note is added is that it gets added to the end of a list of previous notes. The implementation looks like this:

class GenBankEntry:

```
def init (self):
    # . . . other statements . . .
    notes = []
                                                      # initialize the notes field
def get notes(self):
    return self.notes
def add note(self, note):
    self.notes.append(note)
def remove note(self, note):
    self.notes.remove(note)
```

Modification methods are important in another way: with all changes to instances going through these methods, they can control exactly what happens before and after the changes are made. Before making any change, a method could check the validity of the new value(s) supplied. After making the changes, it might update associated information. These are all details that should be hidden behind modification methods.

Defining modification methods also has practical advantages for the class implementer. Such methods can include code to print debugging information, or they can log changes to a file for later examination. They are important places to set debugging breakpoints in order to observe changes being made to instances.

Before moving on, we need to address a subtle problem in the definition of get notes. Because it returns the actual list value of the notes field, external code can modify that list. The same would be true if a dictionary that is the value of a field were returned. Changes to the list or dictionary might require other actions the external code wouldn't know about, such as printing debugging information or updating information located elsewhere. Even if the external code did perform such actions, changes to the representation or required actions would invalidate that code. As always, we want to protect instances from outside meddling.

The solution, like the problem, is a bit subtle. Access methods should never return the actual collection value of a field. Instead, they should return either copies or a generator. We should therefore define get_notes in one of two ways:

```
class GenBankEntry:
    # return copy of list
    def get_notes(self):
        return self.notes[:]

# return generator of the elements of the list
    def get_notes(self):
        return (note for note in self.notes)
```

Action methods

Some classes, like GenBankEntry, are designed simply to *represent* something. Except perhaps for some modification methods and, of course, __init__, they are entirely passive; all of their other methods simply return values.

Other classes are more active; they provide methods that *do* something, in the sense of having an effect outside the class. They may perform input and output operations, create instances of other classes, or perform a wide variety of other sorts of activities. Often, the actions are specific to a particular application.

There isn't much to say about action methods, other than to point out the possibility that a class might have them. They are entirely unconstrained in the arguments they take and the computations they perform. We'll see an example in a parser class later in this chapter.

Support methods

Classes are not limited to defining methods for external code to use. We don't want long method definitions any more than we want long function definitions outside of classes. If a method meant for external use grows too long, it can be divided into multiple methods. As with code extracted from long function definitions, it may turn out that these small methods can be used from several others within the class. Normally they would *not* be used from outside of the class.

We will call these methods "support methods." They should be commented as such to indicate that they are not really part of the class's interface.

Summary

This section examined various kinds of methods in piecemeal fashion. It concludes with a template for a basic class definition. Note that because it plays such a fundamental role in defining instance content, we put __init__ first. Then, since they too are special functions, we put __repr and __str __before the other method groups. Next

come predicates, partly to account for the remaining special method— 1t —and partly because predicates are the simplest methods, just returning a Boolean value.

TEMPLATE

Basic Class

This is a high-level outline of the organization of a typical basic class definition. With the exception of the fundamental methods listed first, there may be any number of methods in each category.

```
class ClassName:
# Fundamental Methods
    def __init__(self, ...):
        # initialize new instance's fields
        # no return
    def repr (self):
        return # string used by the interpreter to print instances
    def str (self):
        return # string used by print and str
# Predicates
    def __lt__(self, other):
        if type(self) != type(other):
             raise Exception(
                 'Incompatible argument to lt : ' +
                str(other))
        return self.somemethod() < other.somemethod()</pre>
    def is some characteristic(self):
        return # True or False
# Access Methods
    def get_something(self):
        return # value obtained by one of:
                    # accessing a field
                    # lookup by key
                    # computation
                    # filtering a collection
                    # searching a collection
# Modification Methods
    def set something(self, ...):
         # change the value of one or more fields based on
         # the parameter values supplied in the call.
         # generally no return value
# Action Methods
    def do something(self, ...):
         # do something that has effects outside the class
```

```
# Private Support Methods
```

```
def helper_method(self, ...):
    # something used by other methods of the class only
```

Class Attributes

Some fields and methods should be associated with the class itself, rather than its individual instances. Typical uses for class attributes are generating a unique ID number for each instance and keeping track of all of a class's instances.

Each class defines its own namespace, just the way modules and functions (including methods) do. That is what allows methods in different classes to have the same name. When a function is defined in a module, its name is added to the module's namespace. When a method is defined in a class, its name is added to the class's namespace.

Actually, "added" is not quite correct: it would be more accurate to say "names and values are added if they're not already there, and otherwise the names are rebound." It is perfectly legitimate, though pointless, to define the same function several times in a row. The first definition adds the function name to its containing namespace, and the others simply rebind what that name means.

Assignment statements also define names, adding (or replacing) names and values within their namespaces. Assignment statements can appear anywhere any other statement can appear. In particular, they can appear inside class definitions alongside the usual method definitions. An assignment statement at the top level of the class definition adds a *class field* to the class's namespace.

Attributes in a class's scope can be accessed in one of two ways:

- As usual, through an instance of the class using dot notation (including self inside one of the class's methods)
- Through the class itself, using dot notation

Class fields

Python classes do not track their instances. That is to say, there is no method you can call on a class to get a list of all its instances. However, we can add instance tracking to GenBankEntry with only two more lines of code: an assignment statement to initialize the class field and a line in the init method to add the new instance to its value. We can use a list or a dictionary, depending on whether we want to be able to look up instances by a key. Since it is very likely that we will want to find instances of GenBankEntry by their GenInfo IDs, we'll use those as the keys in a dictionary.



We use the same capitalization convention for the names of class fields as for the names of classes themselves. This avoids confusion as to which names are for instance fields and which for class fields.

The following code sets up the instance dictionary:

```
class GenBankEntry:
```

```
Instances = {}
                                                              # Initialize the instance dictionary
def init (self):
    \overline{\#} . . . other statements . . .
    self.Instances[self.gid] = self
    # add the new instance to the class's Instances dictionary
```

After creating the GenBankEntry instance from the data we've been using for our examples, we would see one instance in the class's Instances dictionary:

```
>>> GenBankEntry.Instances
                                                         # access to field through class
{'1293613': <GenBankEntry 1293613 U49845.1 'Saccharomyces cerevisiae'>}
```

TEMPLATE

Tracking a Class's Instances

It is often useful for a class to track its instances. All that is required is to assign a name to an empty list or dictionary in a top-level assignment statement and add a line to init to add the new instance to the collection.

```
class UsingList:
   Instances = []
   def __init__(self[, arg, ...]):
        self.Instances.append(self)
class UsingDictionary:
   Instances = {}
   def __init__(self[, arg, ...]):
        self.Instances[self.somefield] = self
```

Another common use for class fields is to keep track of the number of instances that have been generated without tracking the instances themselves. This is useful in order to give each new instance a unique identifier when the class doesn't have a field that can act as one. The identifier would typically be used to define str and/or repr methods that distinguish the instances more helpfully than the awkward default implementation of repr . They can also be used to define 1t if there is nothing else on which to base comparisons for the class.

Why not just track all the instances? Sometimes a program generates many instances of a class during one portion of its code, uses them, then doesn't need them anymore. If they are not tracked, Python can reuse the space they occupy. If they are in a collection in a class, though, Python has no way of knowing if they are still needed. You also might want to track instance counts over multiple executions of a program, which you could do by storing the current count when the program exits and restoring it when the program begins execution later.

Initializing the instance count is easy enough—it's an assignment statement just like the one for tracking the instances. There is one difference, though: there is no way to increment the count that is the value of the class field without using the class's name. Based on the preceding discussion, you might think the following would work:

```
class SimpleClass:
      InstanceCount = 0
   def init (self):
0
          self.number = self.InstanceCount
                                                       # no problem
           self.InstanceCount = self.InstanceCount + 1 # big problem
>>> sc = SimpleClass()
>>> sc.number
                                 # as expected
>>> SimpleClass.InstanceCount
                                 # surprise! why not incremented?
>>> sc.InstanceCount
                                 # mystery!!
```

As you can see, though, it doesn't. Here's what's going on in this code:

- **1** SimpleClass.InstanceCount is bound to **0** when the class is defined.
- 2 The init method binds the field number of the newly created instance to the current value of SimpleClass.InstanceCount, which is 0.
- 3 The full form of the increment statement was used, rather than += 1, to make what happened more explicit. In any assignment statement, the righthand side is evaluated, then the name(s) on the lefthand side is (are) bound to the resulting value(s). The value of self. InstanceCount is still 0, from having been initialized in step 1, so the value of the righthand side is 1. Finally, self. InstanceCount is bound to 1.

The issue lies in the meaning of self.InstanceCount on the lefthand side of the assignment statement. That name refers to a field in the instance, just as self.number does. Python doesn't "understand" that you meant it to refer to the class field. The direct solution to this problem is slightly awkward: use the name of the class explicitly when rebinding the value. We didn't need to do this when modifying a list or dictionary used to track instances, because once the class field was assigned it was never rebound. We'll see a better solution shortly.

Class fields can be used for less dynamic purposes too. There are often names that are specific to the class. These could be assigned at the top level of a module but really should go in the class's namespace. Examples include:

- File paths for input and output
- A format string used in str , repr , and additional output methods
- Minimum and maximum values

One sophisticated use of class fields turns the assignment problem we just looked at into an advantage. As we saw, once a method assigns a field name in an instance, that name refers to the instance's field, not the class's. The value of a name is determined first by looking for a field by that name in the instance, then in its class. This means that a class field can be used as a default value for instances that don't have their own values for a field with that name. Instances that need a different value than the default can assign their own field with that name. The default can even be changed later in the class, and all instances in which the field has not been assigned will take on the new default value. Any instances that did provide a value for the field will keep that value.

Class methods

The benefits of information hiding are important for class fields too. Are they lists, dictionaries, or more complex collections? Maybe their values are instances of other classes the application defines. Changing the contents of a class field's value may require other changes, and as usual, we don't want outside code dealing with these details. We want to be able to define methods to manage class fields the way other methods manage instance fields.

A *class method* is a method for which self is the class itself, not one of its instances. Like a class field, a class method can be accessed using dot notation through either an instance of the class or the class itself.



The syllables that make up the name of a class method, including the first, should be capitalized, as they are in names of classes and class fields.

How are class methods distinguished from instance methods? A special indicator, @classmethod, is placed on the line before a class method definition. (This odd syntax is actually part of a sophisticated aspect of Python that is not discussed further in this book. It is easy enough to use @classmethod without learning anything else about @ notation.) For example:

```
class GenBankEntry:
    Instances = {}
                                                     # initialize the instance dictionary
    @classmethod
                                                     # make the method a class method
    def InstanceCount(self):
        return len(self.Instances)
```

```
@classmethod
         def GetInstances(self):
                                                        # returning generator as discussed earlier
             return (value for value in self.Instance.keys())
         @classmethod
         def Get(self, target):
             """Return the instance whose GID is target"""
             return self.Instances.get(target, None)
Or, if the class keeps its instances in a list instead of a dictionary:
    class GenBankEntry:
         Instances = []
         @classmethod
         def InstanceCount(self):
             return len(self.Instances)
         @classmethod
         def GetInstances(self):
                                                        # returning generator as discussed earlier
             return (value for value in self.Instances)
         @classmethod
         def Get(self, target):
             """Return the instance whose GID is target"""
             for inst in self.GetInstances:
                                                        # using the Find template
                 if target == inst.get gid():
                 return inst
```

Earlier, we encountered problems where an init method incremented a class field used for an instance counter. The solution there was to access the counter through the class name. This is inflexible, though: if the class name is changed, the init method must be changed too. (There is also a more important reason why this is not a good solution, as we'll see later.) A better option is to use a class method. The init method shouldn't be reassigning class fields any more than external code should be reassigning instance fields.

An assignment statement in a class method assigns a name in the class's namespace. Assignment statements in a class definition are executed when the class statement is executed, and assignment statements in a class method are executed when the method is executed. The following template shows how a counter field should really be implemented.

TEMPLATE

Tracking the Instance Count

This template shows how to track instance counts using a class field. It doesn't show what to do with the number field each instance is assigned, but presumably it would be used in the implementation of one or more of _str_, _repr_, and _lt_.

```
class SomeClass:
    InstanceCount = 0
                                                        # initial assignment
    @classmethod
    def IncrementCount(self):
        self.InstanceCount += 1
                                                        # reassignment
    def init (self, args)
        self.number = self.InstanceCount
                                                        # assign an instance field to
                                                        # the value of a class field
        self.IncrementCount()
                                                        # class method call
```

Note that the only variable part of the entire template is the name of the class. (Of course, the number instance field could also be given a different name.) The template's code can be used exactly as shown in any class.

Classes as objects

We've already discussed class fields and class methods with specific examples and explained their use. There is nothing special or mysterious about class fields and methods. Python treats fields and methods of classes and fields and methods of their instances the same way. This is because classes are themselves objects. Remembering this will make it easier to understand how things work.

Class fields and methods are essentially the same as names and functions defined in modules outside of classes. Module functions and values are referenced using dot notation, with the module in front of the dot. Likewise, class fields and methods are referenced using dot notation with the class in front of the dot. A class method can use self before the dot; code outside the class would use the name of the class.

An instance method can call both instance and class methods through self. A field accessed through self in an instance method could be either an instance field or a class field. Python looks first in the instance to see if it contains a field with that name. If it doesn't find it there, it looks in the class. We can't tell by looking at a method call self.methodname() or an expression self.fieldname whether these refer to attributes of the instance or of its class. That is one reason why it is a good idea to use capitalization to distinguish class attribute names from instance attribute names.

An assignment to self. fieldname in an instance method always gives fieldname a value within the instance, whether or not it already has one in the class. Table 5-2 captures all the possibilities.

Table 5-2. Functions and assignments

Internal	Module m Inside m	Class C Inside C	Instance t
Function definition	<pre>def fn():</pre>	@classmethod	<pre>def imeth(self)</pre>
		<pre>def Cmeth(self)</pre>	
Internal call	fn()	self.Cmeth()	self.imeth() or
			self.Cmeth()
Top-level assignment	name = value	Name = value	C.name = value
	Binds m.name	Binds C.Name	
Assignment inside a function fn (like	name = value	self.Name = value	self.name = value
m unless preceded by self, C, or t)	Binds <i>name</i> in <i>fn</i>	binds C. Name	Binds t.name
Scope order (where Python looks)	name	self.Name	self.name
from inside a function (like <i>m</i> unless preceded by self, <i>C</i> , or t)	Function (local)	Class	Instance
preceded by SCIT, e, or e,	Module		Class
	Global		
External access	From outside m	From outside C	
External call	m.name()	C.name()	t.name()
External field	m.name	C.name	t.name

Compare the module and class columns of the table. With the exception of the use of self, there is no difference between the two. Classes and modules are both objects, and their mechanisms are similar. The only difference between the two columns, other than the class's use of self, is that the top-level assignment to a class field does *not* use self self is only a method parameter, not something associated with the class itself. Note that if a name inside a method is not preceded by self, the sequence of scopes in which Python looks for names is the same as if the method had been an ordinary function outside the class.

Comparing the class and instance columns, there are only a few small differences:

- Class method definitions are preceded by @classmethod.
- Methods called through self in an instance method can be either instance or class methods, whereas methods called through self in a class method are always class methods.
- To assign a class field, an instance method must use the class name, exactly as if it were a function outside the class. (It is much better to call a class method that does the assignment so that the name of the class is not part of the method definition.)
- When the value of a field is accessed—not assigned—through self in an instance method, Python looks first in the instance and then in the class.

Class and Method Relationships

There are two fundamental ways that multiple classes can be related to each other. The first implements part/whole relationships among the objects of the classes. The second is as a taxonomy of types. They are both hierarchical relationships, but they differ in their implementation and consequences.

Decomposition

Just as it is important to keep functions focused on relatively narrowly defined tasks, it is important to keep classes focused on relatively narrow representations. When function definitions grow uncomfortably large and perform too many actions, they should be broken up into smaller functions that work together. The same is true for classes: when there is too much information in a class, and—often as a consequence an uncomfortably large number of methods, it is best to break up the class into several that work together.

In addition to being easier to edit and debug, more narrowly focused functions and classes often turn out to be useful beyond their original intended purposes. This happens especially frequently in a field like bioinformatics, where applications often share modules of functions and classes that implement common kinds of data and operations. A class that buries many different kinds of information inside its instances makes it difficult for an application to use only some of that content. In particular, the application might have only some of the data necessary for initializing the instances of the full class.

Dividing up function and class definitions is a programming technique called decomposition. This term refers to the act of breaking up the functions and classes, but it also refers to a design strategy. Decomposing large function and class definitions is good technique. Even better, though, is to think through the likely shape of a part of a program in advance and plan to implement a hierarchy of functions and possibly a group of related classes. Becoming adept at this advanced decomposition takes some practice, so you shouldn't be discouraged if your early efforts in this direction don't seem to accomplish much.

Class decomposition

We know how to divide up a function definition into smaller ones that call each other. But how do we divide up a class? The basic principle is the same one that led to creating a class in the first place: complex structuring collections (as opposed to simple grouping collections) are better represented as instances of a class. We therefore look for fields whose values are complex collections and methods that reach into them. Instances of the new class replace the complex collections in instances of the original class.

Let's take a look at the definition of GenBankEntry. Even though it omits a substantial part of the data found in a GenBank file, it still contains a great deal of information. Its definition would include a large number of access methods that reach into collectionvalued fields to pull out information. The value of features is a list in which each feature is itself a rather elaborate data structure. The situation calls for defining a separate GenBankFeature class and having the value of fields be a list of GenBankFeature instances.

This is an example of turning a collection of complex collections into a collection of instances. The source field is an example of a field whose value is a single complex collection that could be replaced by a single instance. We separated out source from the other features because it plays a different, and more fundamental, role in the GenBankEntry. Since its value has the same structure as the individual features list, it too can be replaced by an instance of GenBankFeature.

A GenBankFeature class would look something like the definition shown in Example 5-2.

Example 5-2. Definition of GenBankFeature

class GenBankFeature:

```
# interesting use of a class field!
   FeatureNameOrder = ('gene', 'promoter', 'RBS', 'CDS')
# Fundamental Methods
   def __init__(self, feature_type, locus, qualifiers):
        self.type = feature type
        self.locus = locus
        self.qualifiers = qualifiers
   def __repr__(self):
         """Return a true __repr__ string"""
        return ('GenBankFeature' +
                 repr((self.type, self.locus, self.qualifiers)))
                # repr of the list of arguments __init__ would take
   def str (self):
        return self.type + '@' + self.locus
# Predicates
   def lt _(self, other):
        if type(self) != type(other):
            raise Exception('Incompatible argument to lt : ' +
                             str(other))
        return self.locus lt(other) and self.type lt(other)
   def is gene(self):
        return 'gene' == self.get type()
   def is cds(self):
        return 'CDS' == self.get type()
```

```
def get type(self):
        return type
    def get locus(self):
        return locus
    def get qualifier(self, name):
        return self.qualifiers.get(name, None)
# Private Support Methods
    def locus lt(self, other):
         """Is this instance's locus "less than" that of other? This could be quite complicated.
         This is just a simple demonstration that only looks for the first integer."""
         assert type(self) == type(other)
                                                 # insisting, not testing
        return ((extract first integer(self.get locus()) or -1) <</pre>
                 (extract_first_integer(other.get_locus()) or -1))
         # extract_first_integer defined elsewhere; not a method
    def type lt(self, other):
        assert type(self) == type(other)
                                                  # insisting, not testing
        return (self.FeatureNameOrder.find(self.get type()) <</pre>
                 other.FeatureNameOrder.find(self.get type()))
```

Note that GenBankFeature.__init__ is defined with individual parameters for each of the values it needs, rather than one collection containing all of them. This is actually a more common approach than just providing a single complex collection argument for the method to disassemble. We'll change GenBankEntry.__init__ similarly, so that we can pass in the GenBankFeature instances after they've been created:

```
class GenBankEntry:
    def __init__(self, accession, gid, source, sequence, features):
        #...
```

Now we define a pair of functions to take the parsed data and create first the GenBankFeature instances and then the GenBankEntry itself:

Separating GenBankFeature from GenBankEntry doesn't mean that we can't define GenBankEntry methods that do things with GenBankFeatures. Such GenBankEntry methods will basically just call the corresponding methods in GenBankFeature. Since the features field in a GenBankEntry instance is a collection of GenBankFeatures, accessing a specific attribute of each of the features would be implemented as a comprehension.

For example:

```
class GenBankEntry:
    def gene names(self):
        return [feature.get qualifier('gene') for feature in features
                if feature.is gene()]
```

This definition doesn't know how to tell if a feature is a gene, so it asks each feature whether it is by using the predicate is gene. Then it collects the names of all features that are genes and returns the resulting collection as its value. When a field's value is not a collection, methods like this can be even simpler. They often just return the result of calling a method on the value of the field. Having replaced source with an instance of GenBankFeature, we can redefine GenBankEntry.get organism as:

```
class GenBankEntry:
    def get organism(self):
        return self.source.get qualifier('organism')
```

Implementing a method primarily by just calling a similar method on a different object is known as delegation.

Method decomposition

Since methods are functions, we can apply the usual functional decomposition strategies to break large methods into smaller ones. Instance methods offer a tactical twist that is sometimes convenient and occasionally powerful: methods can use instance fields to share values among related methods instead of passing them as parameters.

We'll collect the GenBank file parser code into a class, to see how values can be shared as instance fields. While we are rewriting this, we can move the code that creates the GenBankFeature and GenBankEntry instances into the new class, as there is no need for the complex data structure the old code produced or the top-level functions that created instances with them. In doing so, we can compact the code somewhat since we don't need all the features of the original code.

We can also simplify the code by using instance fields to share state among the methods. The code for the parser from Chapter 4 (in "Parsing GenBank Files" on page 148) showed some interesting techniques that are sometimes necessary but that we won't need when implementing the parser as a class. One is that nearly all the functions take an open file as one of their arguments. We can assign an instance field to the open file to be accessed by any method, thereby avoiding having to pass it from one function to the next.

Another situation addressed in that example was that certain functions stopped when they encountered a line meeting some kind of test, but other functions needed that line. As a result, some functions returned the most recently read line, and the functions that called them passed that line as an argument to others. This was one way of "maintaining the state of the process." Instead of passing the line back and forth among the functions, though, we can just use an instance field.

Example 5-3 outlines the structure of a parser class for GenBank files in terms of which methods would call which. A number of predicates that check whether the current line signals the start or end of some section of the file are included. These predicates can be thought of as testing the state of the parser as it moves through the file. They have been italicized in the figure to distinguish them from the more substantial methods. The method read next line is omitted from the outline; it is called from 10 places in the code and simply assigns the instance variable line to the result of a call to readline.

Example 5-3. Using an instance field to share state

```
parse
    get ids
        is at version
    make feature generator
        skip intro
            is at features
        is at sequence start
        read feature
        read qualifiers
            is at feature start
            read qualifier value
                is at attribute start
                is at feature start
        GenBankFeature
    get sequence
        is at sequence end
    GenBankEntry
```

The definition of a GenBankParser class is shown in Example 5-4. The method parse parses the file and returns a GenBankEntry, including its GenBankFeatures. The filename is a parameter to parse, and there are no arguments for init . (An alternative design would be to pass the filename to init and then call parse with no arguments.)

```
Example 5-4. A GenBankEntry parser class
```

```
"""Parse a GenBankEntry file, returning a GenBankEntry instance"""
```

from genbank import GenBankEntry, GenBankFeature

class GenBankParser:

```
def init (self):
    self.line = None
    self.src = None
# interesting use of a class field!
AttributePrefix = (21 * ' ') + '/'
```

```
def is at version(self):
        return self.line and self.line.startswith('VERSION')
   def is at features(self):
        return self.line and self.line.startswith('FEATURES')
   def is at attribute start(self):
        return self.line and self.line.startswith(self.AttributePrefix)
   def is at feature start(self):
        return len(self.line) > 5 and self.line[5] != ' '
   def is at sequence start(self):
        return self.line and self.line.startswith('ORIGIN')
   def is at sequence end(self):
        return self.line and self.line.startswith('//')
# Action
   def parse(self, filename):
        """Use the data in file named filename to create and return an instance of GenBankEntry"""
        with open(filename) as self.src:
            accession, gid = self.get ids()
            feature generator = self.make feature generator()
            source = next(feature generator)
            features = list(feature generator)
                                                       # remainder after source
            seq = self.get sequence()
            return GenBankEntry(accession, gid, source, features, seq)
# Action Support
   def read next line(self):
        """Better to call this than write its one line of code"""
        self.line = self.src.readline()
                                                        # no need to return, just assign
   def get ids(self):
         ""Return the accession and GenInfo IDs"""
        self.read next line()
        while not self.is at version():
            self.read next line()
        parts = self.line.split()
        giparts = parts[2].partition(':')
        return parts[1], giparts[2]
   def skip intro(self):
        """Skip text that appears in self.src before the first feature"""
        self.read next line()
        while not self.is at features():
            self.read next line()
```

```
def get sequence(self):
     """Return sequence at end of file"""
    seq = ''
    self.read next line()
    while not self.is at sequence end():
         seq += self.line[10:-1].replace(' ', '')
        self.read next line()
    return seq
def make feature generator(self):
    """Return a generator that produces instances of GenBankFeature
    using the data found in the features section of the file"""
    self.skip intro()
    while not self.is_at_sequence_start():
        yield self.read feature()
def read feature(self):
    """Return an instance of GenBankFeature created
    from the data in the file for a single feature"""
    feature type, feature locus = self.line.split()
    self.read next line()
    return GenBankFeature(feature type,
                            feature locus,
                            self.read_qualifiers())
def read qualifiers(self):
    qualifiers = {}
    while not self.is at feature start():
        parts = self.line.strip()[1:].split('=')
        key = parts[0]
        value = '' if len(parts) < 2 else parts[1]</pre>
        if value and value[0] == '"':
             value = value[1:]
                                                    # remove first quote; last removed later
        qualifiers[key] = self.read qualifier value(value)
    return qualifiers
def read qualifier value(self, value):
    """With value, the string after a qualifier name, and its equal and quote, keep
    reading lines and removing leading and trailing whitespace, adding to
    value until the final quote is read, then return the accumulated value"""
    self.read next line()
    while (not self.is at attribute start() and
            not self.is at feature start()):
        value += self.line.strip()
         self.read next line()
    if value and value[-1] == '"':
        value = value[:-1]
                                                    # remove final quote
    return value
```



It isn't actually necessary for init to initialize file and line, the fields that maintain the state of the parsing process. Fields can be assigned for the first time in methods other than init: until the first assignment of the field is executed, wherever it occurs, the instance just doesn't have a field by that name. However, readers of the code will expect to see all fields initialized in init, and are unlikely to want to read through the entire class's code checking for field assignments. Therefore, it is good practice to assign all fields the instance will use in the init method, even if only to None. In essence this is a form of documentation, identifying which fields are used by the class's methods.

Even if a method is called from only a few others, it can be useful to assign a field to its arguments. This can be done in the method itself, or it can be done by a calling method before a call, in which case the values will not be included in the call. In fact, even if a method assigns names of its own for local use it can be helpful to turn those into field assignments.

An additional benefit of using this technique is that it makes debugging much easier. The field assignments capture the state of the instance. After the program calls a method on the instance, or while the program is stopped in the debugger, you can look at the values of its fields to figure out what has happened. (The Python debugger is introduced at the end of the next chapter.) This can be easier than tracing or stepping through a series of method calls to watch what values parameters take on.

Singleton classes

Classes that define parsing actions for a particular file format are a common application of object-oriented programming techniques. In this case, the motivations for the definition of a class to parse GenBank entry files were purely technical. Reconfiguring the previous code as a set of methods made certain aspects of the code easier to read, write, and debug, and using instance fields to share state among methods avoided some of the complexity of the functional solution.

To use GenBankParser, code would create an instance and call its parse method, giving it the name of the file containing the GenBank entry text. That method returns an instance of GenBankEntry. Suppose we want to write an application that parses multiple GenBank files. Do we need a new instance of GenBankParser for each file? There are two things we need to know to answer that question:

- After parse returns, is there any state left in the instance that we'll want to retrieve
- After parse returns, is there any state left in the instance that would get in the way of using it again?

If the answer to both questions is "no," as it is in this case, we can reuse the same instance to parse multiple files. A different reason for needing only one instance of a

class is that it represents the application itself, the state of the user interface, or something else that must exist during the entire execution of the application. This is different from reuse: it is continual use.

Classes of which an application needs only one instance are called *singleton classes*. It may seem strange to create a class that has only one instance: most classes are defined in order to have many instances, each with their own state. There's nothing wrong with it, though, and singleton classes are rather common. The concept is introduced here so you'll recognize such classes for what they are.

One unusual thing about singleton classes is that it doesn't matter whether their implementations use all instance fields and methods or all class fields and methods. If we were to copy the definition of GenBankParser, delete the init method, and put @classmethod in front of every method, it would work exactly the same as the original definition. The only difference is that instead of creating an instance and calling parse on it, we would just call GenBankParser(filename) directly.

Sometimes there isn't much reason to choose one approach over another. If there are many methods that might be called from outside the class (instead of just one, as in this example), it is somewhat easier to call them using the name of the class than through a name bound to a single instance of the class. That is because once the name of the class is imported into a module's namespace, it can be used directly, which is much easier than passing an instance among many functions as one of their arguments.

Inheritance

It often happens that you want to design and implement a class that is a lot like another. You could copy the code from the other class to use as a starting point and then modify it slightly, but that strategy has many weaknesses:

- The more copies of a body of code there are, the more work it takes to maintain them.
- The more copies of code there are that need changing, the more likely it is that mistakes will be made in making the changes.
- The larger the code grows, the more there is to read and understand.
- The larger the code grows, the more effort it takes to navigate through and edit the code.

Object-oriented languages provide a powerful mechanism called *inheritance* that allows you to define a new class that differs from an old one only in the methods it defines. This new class is called a *subclass*, and the class on which it is based is called a *super*class. (The terms base class and derived class are also sometimes used.) When a class is defined with one or more superclasses, its definition implicitly includes all the methods and class fields of each of its superclasses. Those, in turn, contain all the methods and class fields of their superclasses. Before it defines anything at all, the new subclass inherits all of the methods and class fields of the entire chain of superclasses.

Defining subclasses

A subclass is defined by following its name with the name of its superclass, enclosed in parentheses, as shown in Example 5-5.

Example 5-5. Defining a subclass

```
class BaseSequence:
   def init (self, seqstring):
        self.seg = segstring
   def get sequence(self):
        return self.seq
   def gc content(self):
         """Return the percentage of G and C characters in base_seq"""
        seq = self.get sequence.upper()
        return (seq.count('G') + seq.count('C')) / len(seq)
```

class RNASequence(BaseSequence): # subclass # . . .



The parentheses can actually contain a list of superclass names. Having more than one superclass is called multiple inheritance. Multiple inheritance can be used for a wide range of purposes, from specific implementation techniques to high-level architectural mechanisms. Multiple inheritance is not all that difficult to understand or use in Python, but it leads to complexities and issues beyond the scope of this book.

A method in a class can refer to its superclass by calling the built-in Python function super. The word self does not appear either before super or within its argument list. The call to the function super takes the place of self in a method call. The result of super is an object that represents the same instance as self, but for which methods are obtained from the superclass.



Python 2: In Python 2 code you may occasionally see a class defined with object as its superclass. In Python 3 object is a superclass of every class, so class definitions do not include it explicitly.

We need super as soon as we start defining subclasses, because the init method of a subclass *must* call the init method of its superclass. This is so that each class can initialize the fields of the new object according to its own init procedure. Example 5-6 shows a small example.

class RNASequence(BaseSequence):

```
CodonTable = {'UUU': 'F', 'UCU': 'S', 'UAU': 'Y', 'UGU': 'C',
              # . . .
              }
@classmethod
def translate codon(self, codon):
    return self.CodonTable[codon.upper()]
def init (self, seqstring):
    super(). init (seqstring) # note that super does not use self
def translate(self, frame=1):
    """Produce a protein sequence by translating this
    RNA sequence starting at frame 1, 2, or 3""
    return ''.join([self.translate codon(self.get sequence()[n:n+3])
                     for n in
                     range(frame-1,
                            # ignore 1 or 2 bases after last triple
                           len(self.get sequence()) -
                           (len(self.get_sequence()) - (frame-1)) % 3,
                           3)])
```

This example defines the class BaseSequence, which simply stores a sequence (presumably a string) and provides an access method for it. It also defines RNASequence as a subclass of BaseSequence. Because BaseSequence defines get_sequence, that method can be called on instances of RNASequence. The translate method can be called on an instance of RNASequence, but not on an instance of the more general BaseSequence, since it doesn't define that method. Although an RNASequence has no initialization work to do for its own purposes, it must forward to BaseSequence.__init__ the arguments it expects:

Initialization functions often check the validity of their arguments. We'll extend the definition of RNASequence.__init__ to use a class method that raises an exception if there is an invalid character in seq:

```
@classmethod
def invalid_chars(self, seqstring):
    return {char for char in seq if char not in 'UCAGucag'}

def __init__(self, seqstring):
    invalid = self.invalid_chars(seqstring)
```

```
if invalid:
    raise Exception(
        'Sequence contains one or more invalid character]: ' +
        str(invalid))
super().__init (seqstring)
```

Next, we'll add a DNASequence class with a very slightly different definition of isinvalid. Instead of translate, it includes a transcribe method that returns its RNA complement:

```
class DNASequence(BaseSequence)
    # construct a string translation table for use with str.translate
    TranslationTable = str.maketrans('TCAGtcag', 'AGUCaguc')
    @classmethod
    def invalid chars(self, seqstring):
        return {char for char in seq if char not in 'TCAGtcag'}
    def __init_ (self, seqstring):
        invalid = self.invalid chars(seqstring)
        if invalid:
            raise Exception(
                 'Sequence contains one or more invalid characters: ' +
                 str(invalid))
        super().__init (seqstring)
    def transcribe(self):
        """Produce an instance of RNASequence that is
        the transcribed complement of this sequence"""
        return RNASequence(
                                 # use string translation table to transcribe
             self.get sequence().translate(self.TranslationTable))
```

Classes related by inheritance form a *class hierarchy*. Conceptually, this is a taxonomy of types, analogous to a biological taxonomy. However, the relationship is more than just conceptual, due to the technical consequences of the superclass/subclass relationship. The (simple) taxonomy for the preceding example is shown in Figure 5-1.

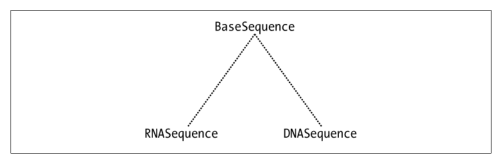


Figure 5-1. A class hierarchy

Factoring out common code

class BaseSequence:

Notice the repetition of code in RNASequence and DNASequence in the steps taken to validate the seq argument of init . The only difference is two characters in the string of invalid chars. Otherwise, invalid chars and init are exactly the same in the two classes. Were we to notice such duplication in an ordinary function, we would pull out the common code and put it in a new function, calling it from the old ones. Slight differences in the old code would be handled by adding parameters to the new function.

With classes, we can do even better than that: methods that are duplicated in sibling classes can be moved up to a common superclass. Removing such duplication is one of the most common uses of inheritance. By analogy with the similar notion from algebra, this reorganization is often called *factoring out*.

Example 5-7 shows revised—and shorter—definitions of the classes. The method invalid chars has been moved to BaseSequence, and the test for invalid characters has been moved from RNASequence. init and DNASequence. init into BaseSequence init .

Example 5-7. Factoring out common code to a superclass

```
@classmethod
    def invalid chars(self, seqstring):
        return {char for char in seq if char not in self.ValidChars}
    def __init__(self, seqstring):
        invalid = self.invalid chars(seqstring)
        if invalid:
            raise Exception(
                type(self).__name__ +
                'Sequence contains one or more invalid characters: ' +
                str(invalid))
        self. seq = seqstring
    def get sequence(self):
        return self. seq
    def gc content(self):
         """Return the percentage of G and C characters in the sequence"""
        seq = self.get sequence.upper()
        return (seq.count('G') + seq.count('C')) / len(seq)
class RNASequence(BaseSequence):
    ValidChars = 'UCAGucag'
    CodonTable = {'UUU': 'F', 'UCU': 'S', 'UAU': 'Y', 'UGU': 'C',
                   # . . .
                   }
    @classmethod
    def translate codon(self, codon):
```

```
return self.CodonTable[codon.upper()]
    def init (self, seqstring):
        super(). init (seqstring)
    def translate(self, frame=1):
        """Produce a protein sequence by translating this
        RNA sequence starting at frame 1, 2, or 3"""
        return ''.join([self.translate codon(self.get sequence()[n:n+3])
                         for n in
                         range(frame-1,
                                # ignore 1 or 2 bases after last triple
                                len(self.get sequence()) -
                                (len(self.get sequence()) - (frame-1)) % 3,
                                3)])
class DNASequence(BaseSequence):
    ValidChars = 'TCAGtcag'
    # construct a string translation table for use with str.translate
    TranslationTable = str.maketrans('TCAGtcag', 'AGUCaguc')
    def __init__(self, seqstring):
        super(). init (seqstring)
                                                                  # note absence of self
    def transcribe(self):
        """Produce an instance of RNASequence that is
        the transcribed complement of this sequence"""
        return RNASequence(
            self.get sequence().translate(self.TranslationTable))
```

To generalize the definition of invalid chars, the strings the old definitions contained have been replaced by a class variable, ValidChars. When the method is executed, self will be an instance of either RNASequence or DNASequence, so in evaluating the expression self. ValidChars Python will look first in the instance, then in the instance's class, where it will find the value. (If the instance's class did not define the class field, Python would next look for it in the superclass.) Example 5-8 demonstrates this process in the creation of a new DNASequence.

```
Example 5-8. Referencing a subclass class value
```

```
DNASequence('ATTCTGGC')
   → DNASequence. init ('ATTCTGGC')
        → BaseSequence.__init__('ATTCTGGC')
            → BaseSequence.isinvalid('ATTCTGGC')
                    which references self.ValidChars
                         1. does self have a ValidChars field? no
                         2. does self's class have a ValidChars field? yes
                         [3. if it did not, the superclass of self's class
                              would be examined for a ValidChars field]
```

Generalization

It is often the case that after defining two apparently independent classes, some commonality becomes apparent. Such situations motivate the definition of a new class. This new class is a generalization of its subclasses. In truth, any class can be described as a generalization of its subclasses, but the term specifically calls attention to the action of creating a new superclass to capture code common to some existing classes.

As an example of generalization, suppose we want to add ProteinSequence to our group of classes. It will share some, but not all, of BaseSequence's implementation. Therefore, we'll define a more generalized class called Sequence. This leads to the following changes:

- BaseSequence is changed to inherit from Sequence.
- A new class, ProteinSequence, is defined with Sequence as its superclass.
- The initialization of seq is moved to Sequence. init .
- The method sequence is moved to Sequence.
- The method gc content stays in BaseSequence, because it is only applicable to base sequences, not to amino acid sequences.
- The method valid chars is moved to Sequence.

While we're at it, we can also add a new class called AmbiguousDNASequence, which allows sequences to have ambiguity codes in addition to DNA symbols. At first sight it might look like this should be a subclass of DNASequence. However, ambiguous sequences cannot be transcribed into RNASequences, and it is rarely appropriate to define a subclass when its superclass has methods that are inapplicable to the new class's instances. Thus, we'll make Ambiguous DNA Sequence a direct subclass of Base Sequence.

Now that we have a ProteinSequence class we can change RNASequence.translate to return an instance of that instead of a string, the way DNASequence returns an instance of RNASequence. We'll also define a str method for Sequence that simply returns the sequence. Because these classes are all instantiated with a single string argument, we can define an appropriate repr as well.

For both ProteinSequence and AmbiguousDNASequence, we'll assign appropriate ValidChars values and define the usual __init__ methods. We'll also give ProteinSequence a long string method that returns a string with the one-letter codes replaced by the corresponding three-letter ones. We'll use another class field to define a dictionary to implement that and create a list of valid characters by extracting its keys. Note that during the execution of the class definition statement, class fields are just ordinary names. One assignment can use the result of a previous assignment. There is no self at that point, because the class is not defined until the class statement has completely finished executing.

Example 5-9 shows the definitions of the new classes and the changes to the old ones.

```
class Sequence:
```

```
@classmethod
    def invalid chars(self, seqstring):
         return {char for char in seq if char not in self.ValidChars}
    def init (self, seqstring):
         invalid = self.invalid chars(seqstring)
         if invalid:
              raise Exception(
                   type(self).__name__ +
                    " contains one or more invalid characters: " +
                   str(invalid))
         self. seq = seqstring
    def get sequence(self):
         return self. seq
class BaseSequence(Sequence):
    def init (self, seqstring):
         super(). init (seqstring)
    def gc content(self):
           """Return the percentage of G and C characters in the sequence"""
         seq = self.get sequence.upper()
         return (seq.count('G') + seq.count('C')) / len(seq)
class ProteinSequence(Sequence):
    ThreeLetterCodes = \{ \text{ # including ambiguity codes B, X, and Y } \}
                   'A': 'Ala', 'B': 'Asx', 'C': 'Cys', 'D': 'Asp', 'E': 'Glu', 'F': 'Phe', 'G': 'Gly', 'H': 'His', 'I': 'Ile', 'K': 'Lys', 'L': 'Leu', 'M': 'Met', 'N': 'Asn', 'P': 'Pro', 'Q': 'Gln', 'R': 'Arg', 'S': 'Ser', 'T': 'Thr', 'V': 'Val', 'W': 'Trp', 'X': 'Xxx', 'Y': 'Tyr', 'Z': 'Glx'}
    ValidChars = ''.join(ThreeLetterCodes.keys())
    ValidChars += ValidChars.lower()
    TranslationTable = str.maketrans(ThreeLetterCodes)
    def long str(self):
         return self.get sequence().translate(self.TranslationTable)
         # or ''.join([ThreeLetterCodes[key]
                            for key in self.get sequence()])
    def init (self, seqstring):
         super(). init (seqstring)
class RNASequence(BaseSequence):
    ValidChars = 'ucagUCAG'
```

```
CodonTable = {'UUU': 'F', 'UCU': 'S', 'UAU': 'Y', 'UGU': 'C',
                   }
   @classmethod
   def translate codon(self, codon):
       return self.CodonTable[codon.upper()]
   def init (self, seqstring):
        super().__init__(seqstring)
   def translate(self, frame=1):
        return ProteinSequence(
            ''.join((self.translate codon(self.get sequence()[n:n+3])
                     for n in
                     range(frame-1,
                           len(self.get sequence()) -
                            ((len(self.get sequence()) - (frame-1)) % 3),
                           3))))
class DNASequence(BaseSequence):
   ValidChars = 'TCAGtcag'
   # construct a string translation table for use with str.translate
   TranslationTable = str.maketrans('TCAGtcag', 'AGUCaguc')
   def __init__(self, seqstring):
        super(). init (seqstring) # note absence of self
   def transcribe(self):
        """Produce an instance of RNASequence that is
        the transcribed complement of this sequence"""
        return RNASequence(
            self.get sequence().translate(self.TranslationTable))
class AmbiguousDNASequence(DNASequence):
   ValidChars = 'TCAGBDHKMNSRWVY'
   ValidChars += ValidChars.lower()
   def __init__(self, seqstring):
        super(). init (seqstring)
```

The inheritance relationships among these classes are shown in class hierarchy diagrammed in Figure 5-2.

The changes made in Example 5-9 left BaseSequence empty. Let's add a method that applies to all kinds of BaseSequences, but not to ProteinSequences: complement. We'll use translation tables to implement it, as we did for DNASequence.transcribe. We'll even take the somewhat unusual step of defining complements for ambiguity codes:

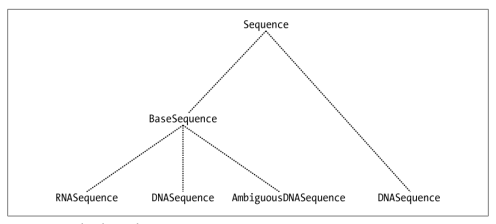


Figure 5-2. A class hierarchy

```
class BaseSequence(Sequence):
    # . . .
    def complement(self):
        return self(self.get sequence().translate(self.ComplementTable))
    def gc content(self):
         """Return the percentage of G and C characters in the sequence"""
        seq = self.get sequence.upper()
        return (seq.count('G') + seq.count('C')) / len(seq)
class RNASequence(BaseSequence):
    ComplementTrans = str.maketrans('UCAGucag', 'AGUCaguc')
    # . . .
class DNASequence(BaseSequence):
    ComplementTable = str.maketrans('TCAGtcag', 'AGTCagtc')
    # . . .
class AmbiguousDNASequence(DNASequence):
    ComplementTable = str.maketrans('TCAGRYMKSWNBVDH', 'AGTCYRKMSWNVBHD')
```

Now we can invoke complement on instances of any subclass of BaseSequence, getting results appropriate to each type of instance.

Subclass methods

Methods defined in a subclass can serve a number of different purposes relative to the methods of their superclasses:

Addition

The subclass extends the superclass's repertoire by defining new methods.

Compounding

A subclass method can invoke several superclass methods. This has the effect of combining the actions of those methods.

Blocking

Very occasionally, especially in elaborate inheritance hierarchies, it makes sense to define a class as a subclass of another even if it does not support a few of the superclass's methods. For instance, a simplistic view of real-world biology would say that a penguin is a kind of (i.e., subclass of) bird, except that it doesn't fly. Usually, though, it is better to refactor (i.e., rearrange) the class hierarchy to eliminate this conflict. We did this when adding ProteinSequence to our earlier example.

You have three options when defining a method that blocks an inherited one: do nothing, print or log a "not supported" warning, or raise an exception.

Extension

The subclass method calls the superclass method before, in the middle of, or after performing some other action(s). Each class's init method is necessarily an extension of its superclass's init method. For other methods, extension can take many forms. The subclass can add additional verification of values used by the superclass's methods, add printing or logging facilities, check security restrictions, store or update information beyond what is managed by the superclass method, and so on.



An extension method's call to the corresponding superclass method is not limited to passing the values of its parameters. There are many kinds of opportunities for passing values that have been changed or replaced in some way.

Definitions of init are usually more constrained. They almost always begin with a call to super.__init__, with an unmodified set of arguments corresponding to the inherited parameter list. If the subclass's init defines additional parameters, those will typically be processed after the call to the superclass's init .

Overriding

There is nothing problematic about defining a method in a subclass that simply replaces one with the same name in its superclass. This is called *overriding*. Unlike extension methods, a method that overrides one its superclass defines does not use super. Normally, the subclass method serves the same purpose as the superclass method; the only difference is in its implementation.

One method that needs overriding in our example is gc content. In computing the proportion of G and C bases, we divided the total count of the two by the length of the sequence. However, assuming it makes sense to compute gc content for DNA sequences that include ambiguity codes, we need a different definition for gc content in AmbiguousDNASequence. In computing the sum of the G and C bases

it should also include occurrences of the ambiguity code S, since that stands for either a G or a C. It should then compare the sum of those counts not to the entire length of the sequence, but rather to the count of symbols that are either definitely G or C (which includes S) or definitely A or T (which includes W). For the purposes of this computation, we must ignore other ambiguity codes. The definition, a typical example of overriding, would be:

class AmbiguousDNASequence(DNASequence):

```
def gc content(self):
       "Return the percentage of G and C characters in the sequence, counting
    only bases that are either definitely G or C or definitely A or T""
    seq = self.get sequence.upper()
    gc count = (seq.count('G') + seq.count('C') + seq.count('S')
    at count = (seq.count('A') + seq.count('T') + seq.count('W')
    return gc count / (gc count + at count)
# . . .
```



Overriding and extension are different from the other techniques in a very important respect: because the subclass method replaces the superclass method in calls through instances of the subclass, its parameter list should conform to that of the superclass method. That is, it should look like the superclass method so that it can be used without distinguishing the type of the instance through which it is called. This means that the subclass's parameter list should begin with the same parameters as the ones in the superclass method's parameter list.

Other arguments may be added after those for the benefit of code that knows it is dealing with instances of the subclass. Such arguments must be optional so that other code does not need to supply values for them; they must be defined as keyword arguments that follow the parameters of the superclass method.

Tips, Traps, and Tracebacks

Tips

- Don't capitalize the names of class files: Python filenames should be all lowercase. Confusion can result from having modules and classes with the same names. Capitalizing all your class names and leaving filenames in lowercase avoids problems.
- The class definitions in this book's examples generally use the same name for the parameters of init methods as the fields that get initialized with their values. This is a stylistic preference, not a requirement.
- Most methods of a class should use its own get and set methods to access its fields instead of accessing them directly. The advantages of information hiding are

important for the class developer too, not just for other programmers who use the class. It may not seem important when you first develop a class, but when you come back to it later, it is easy to forget what the fields contain. Also, many get methods may compute, search for, or look up values that aren't directly stored in a field. Dictionary-valued fields are particularly problematic in this regard:

- Quite often, other methods in the class will do something with the dictionary's entries. They should call get entries. Otherwise, when the developer comes back to the class after a while and sees a field with the name entries, he may well try to use it directly, thinking it is a sequence. The errors that result when keys are used instead of values are distracting at minimum, and often quite difficult to decipher.
- —A very common mistake with dictionary-valued fields is using them directly in an iteration, comprehension, or generator expression without calling keys, values, or items. Doing so is the same as calling keys, which is not always what you want. An access function get entries that accesses the values of a dictionary-valued field would normally be coded as a generator expression:

```
def get entries(self):
   return (entry for entry in self.entries.values())
```

- -Calling the field entry dict might avoid that particular confusion, but there is still the problem that if the representation is changed from a list to a dictionary, or the other way around, the name of the field will have to be changed throughout the class. More importantly, the code that used to use entry dict.values will have to be changed to use entries directly. This can be avoided by using get entries in the class's own methods. Then, a change in the representation requires changing only get entries (and perhaps one or more "set" methods).
- When extending an inherited method, think carefully about whether you want the superclass's method to be called before, in the middle of, or after the body of the subclass's method. For init, the call to super should almost always be first. Otherwise, it depends on the way the methods are designed to work, especially if the superclass method calls others.

Information Hiding

We have talked extensively about defining a functional interface to classes and instances and insisting that external code use it. Yet the mechanisms and techniques shown in this chapter are insufficient to prevent external code from directly accessing class and instance fields or calling methods meant to be used only from other methods of the class. Information hiding can be enforced in Python at two different levels:

 Prefixing a method or field with a single underscore is a convention indicating that those names should not be used outside the class, except perhaps by closely related classes. It has no effect, but it does highlight the class implementer's intention that the method or field should not be used outside the class.

Prefixing a name with two underscores makes that name unavailable outside the class; this restriction is implemented by Python. For example, if an init method assigned self. name, external code would not be able to access that field —it would be forced to use the get name method the class presumably supplies.

Do not *end* a name with two underscores—this is a different convention than Python's special methods, which both begin and end with two underscores.

Traps

- Omitting self from the parameter list of a method is a common mistake. The error that results is usually a complaint about the method being called with one too many arguments.
- Omitting self. before a reference to a field or method of the same class inside of a method is another common mistake; this will typically manifest itself as a name error, since the field or method name is not likely to have a binding outside of the instance's namespace.
- If an instance is lacking a field you expected to find, make sure the assignments in init begin with self. Otherwise, the names assigned will just be local to the method's namespace, and they'll disappear when the method exits.
- No assignments at the top level of a class, including assignment of default values for method parameters, may refer to self. The class does not exist until its definition statement has completed execution. At that point Python attaches its namespace to it. The same thing happens with modules: a module being imported does not get created until all of its statements have been executed, at which point Python creates a module object and attaches to it the namespace containing its definitions.
- In an instance method, reassignment to a class field through self assigns that field in the instance, not the class field as intended. You must use the name of the class to rebind the class field from within an instance method: for example, GenBankEntry.count instead of this.count. You can access the value of the class field through self, but you cannot rebind it that way. Assigning a name through self binds a field in that instance the way assigning a name inside a function definition binds that as a name to the function.
- When you run Python from the command line or run a program using IDLE's Run menu command (F5), your classes are loaded into a clean Python. If, however, you are using Python interactively outside of IDLE, you must understand the effects of executing and importing code. This is much more of an issue when using classes, and even more so with multiple modules:
 - —Reimporting a module has no effect: Python keeps a list of modules that have been imported and doesn't import ones that are on that list. There are ways to "reload" modules, but they are not meant for casual use. If you are not using IDLE (or another IDE that clears the status of the Python environment when a

- file executes), and you are executing the code in one file that imports code from another file that has changed, you pretty much have to restart Python.
- Trouble can arise with old definitions even if you have all your code in one file. When you reexecute a class's definition, the old instances continue to refer to the old class definition, with its old method definitions. A module that creates instances of another module's classes may also end up referring to the old classes. This is a very confusing area of Python use. While inconvenient in some respects, IDLE's automatic restart eliminates this problem too. If you are getting mysterious results after editing and reexecuting class definitions outside of IDLE, try restarting Python.

Tracebacks

Here are some representative error messages:

```
AttributeError: type object 'super' has no attribute 'draw'
    super was used without parentheses.
```

```
NameError: global name 'attribute' is not defined
```

The name attribute (a field or method) was not preceded by self. inside a method.

```
TypeError: method() takes no arguments (1 given)
    A method's parameter list did not include self.
```

```
TypeError: unorderable types: Classname() < Classname()</pre>
```

No lt method was defined for Classname, and sorted was called on a sequence of its instances or list.sort on a list of its instances.

The following does not cause a traceback, but it is probably not what you expected when you tried to call a method such as get_gid while omitting the parentheses:

```
>>> gb
 <GenBankEntry 1293613 U49845.1 'Saccharomyces cerevisiae'> # uses __repr__
>>> gb.get gid
<bound method GenBankEntry.get gid of # line break inserted manually for formatting purposes</pre>
 <GenBankEntry 1293613 U49845.1 'Saccharomyces cerevisiae'>>
```

This is simply another form of leaving out the parentheses in an expression intended to be a function call. For instance:

```
>>> print
<built-in function print>
```

A bound method is a method associated with (bound to) a particular instance. Before it is bound, it is just a function like any other:

```
>>> GenBankEntry.get gid
<function get gid at 0x1c7d228>
```

Once it is called through an instance, however, the method's type changes to bound method and its printout includes the instance to which it is bound.

Utilities

This chapter surveys many Python modules that are particularly important for bioinformatics programming. We will look at utilities that help you connect Python to its external environment, deal with filesystems, work with text, and debug your programs, among others. These aren't particularly exciting, but it's easy enough to learn how to use them. You'll need some of them to deal with operating systems, filesystems, dates and times, and so on. Others will likewise prove extremely useful in the right circumstances, giving you ways to do in a few lines of code things that could take pages otherwise. Some of the modules aren't necessary for simple scripts or initial development but provide the features users expect from mature programs.

System Environment

The first set of modules we'll look at are those that allow Python to interact with its external environment. The categories included here are dates and times, command-line processing of various kinds, email, and logging.

Dates and Times: datetime

The datetime module defines classes that represent a date, a time, a combination of date and time, and a couple of others. There are some basic ways of manipulating instances of these classes, as well as some more elaborate ones that we won't consider here.

Classes

The datetime module defines five classes. Their representations are simple, consisting only of several fields with primitive types. This allows them to define <code>__repr__</code> methods that print exactly what you would enter to create one. You'll see such output when you enter into the interpreter an expression whose value is an instance of one of these classes. They also define <code>__str__</code> methods for the more human-oriented output used by print. The five classes are:

```
datetime.date
```

Represents a date with attributes year, month, and day

```
datetime.time
```

Represents a time with attributes hour, minute, second, microsecond, and tzinfo (time zone information)

datetime.datetime

Represents a combination of date and time with the attributes of each

```
datetime.timedelta
```

Represents the difference between two dates, two times, or two datetimes, with attributes days, seconds, and microseconds

```
datetime.tzinfo
```

Represents time zone information

Note that unlike many classes, these don't define "access" methods to get the values of their attributes. Instead the attributes are accessed directly using dot notation, e.g., date1.year. It is very important to note that as with strings, instances of the classes in this module are immutable. There are methods to create one instance based on the values of another's attributes, but once an instance is created its attribute values cannot be changed.

Instance creation

Like any other type, the types in the datetime module can be called as functions to create new instances:

```
datetime.date(year, month, day)
datetime.time(hour[, minute[, second[, microsecond[, tzinfo]]]])
datetime.datetime(year, month, day[, hour[, minute[, second[, microsecond[,
tzinfo]]]]])
datetime.timedelta([days[, seconds[, microseconds[, milliseconds[, minutes[,
hours[, weeks]]]]]]])
```

The classes also provide methods to create instances based on the current date and/or time:

```
datetime.date.today()
datetime.datetime.now([tz])
datetime.datetime.combine(date, time)
```

Return a datetime with the date portion taken from date and the time portion taken from time

There are also methods that create new instances from existing ones, possibly of a different class:

date1.replace(year, month, day)

Returns a new date with the same attribute values as date1, except for those for which new values are specified (usually as keyword arguments)

```
time1.replace([hour[, minute[, second[, microsecond[, tzinfo]]]]])
```

Returns a new time with the same attribute values as time1, except for those given new values by whichever keyword arguments are specified

datetime1.date()

Returns a new date with the day, month, and year of datetime1

```
datetime1.time()
```

Returns a new time with the hour, minute, second, and microsecond of datetime1 datetime1.replace([year[, month[, day[, hour[, minute[, second[, microsecond[, tzinfo]]]]]]])

Returns a new datetime with the same attribute values as datetime1, except for those given new values by whichever keyword arguments are specified

Operations

Instances of these classes support a few arithmetic and comparison operations. Table 6-1 shows the arithmetic operations supported by instances of date.

Table 6-1. Operations supported by date instances

Operation	Result
date1+time delta1	date1+timedelta1.days
date1 - time delta1	date1 - timedelta1.days
date1 - date2	Atimedelta
date1 < date2	True if date1 is earlier than date2

The same operations are supported by instances of datetime with equivalent interpretations. However, the only operation supported by time instances is <. Table 6-2 shows operations that can be performed with instances of timedelta.

Table 6-2. Operations supported by timedelta instances

Operation	Result
t1+t2	Sum
t1 - t2	Difference
t1*i,i*t1	Product of $t1$ and integer i
t1//i	"Floor" division, as with numbers
-t1	A new timedelta whose attribute values are the negation of t1's

Operation	Result
abs(t1)	A copy of $t1$, except that a negative value of days is made positive
comparisons	The usual six: ==, !=, <, <=, >, >=

Methods

In addition to being able to access the attributes described in the preceding sections using the usual dot notation, the following methods are often useful (the module's classes define many more, but most are used primarily in low-level system development):

```
date1.isoweekday()
    Returns the day of the week of date1 as an integer, with Monday = 1
date1.isocalendar()
    Returns a tuple of date1's values with the following elements:
        (ISO year, ISO week number, ISO weekday)
date1.isoformat([sep])
    Returns a string representing date1 in the standard format:
        YYYY-MM-DD
time1.isoformat([sep])
    Returns a string representing time1 in the standard format:
        HH:MM:SS.mmmmmm
    omitting the microseconds if 0
datetime1.isoweekday()
    Returns the day of the week of datetime1 as an integer, with Monday = 1
datetime1.isocalendar()
    Returns a tuple of datetime1's values with the following elements:
        (ISO year, ISO week number, ISO weekday)
datetime1.isoformat([sepchar])
    Returns a string representing datetime1 in the standard format:
        YYYY-MM-DDsepcharHH:MM:SS.mmmmmm
```

System Information

When a program is started, it may be given command-line arguments that are in effect parameters of the program itself. While it is executing, it can read from and write to

separates the date and time portions of the string ('T', if omitted)

omitting the microseconds if 0; sepchar is an optional one-character string that

predefined standard streams that connect it to the environment in which it is running. These fundamental features, along with access to details about Python's internal environment, are implemented by the sys module. Programs may also occasionally need to access aspects of the external environment, including the computer's internal clock and information about other time-related details such as the time zone and daylight savings time settings. The time module provides facilities for that.

The Python runtime environment: sys

The sys module provides access to a great deal of technical information about the Python implementation. It provides mechanisms for modifying various aspects of the interpreter, such as the prompt. Following are frequently used values defined by the module (remember, you must use either import sys or from sys import name for whatever name you want to use):

sys.argv

This is the single most important name defined in sys. When Python is run from the command line, sys.argv is a list of strings containing the parts of the command line. The first—sys.argv[0]—is the name of the Python program being run. You will use this often. Consider the following very brief demonstration program:

```
# File showargs.pv
import sys
for arg in sys.argv:
    print(arg, end=' ')
```

Running this program from the command line simply prints the name of the file and the command-line arguments:

```
% python showargs.py a 1
showargs.py a 1
```



If your program expects a certain number of arguments, you should always check that no fewer arguments than expected and no more than allowed have been provided. This should be done at the end of the file, inside a test that determines whether name is ' main ', since you wouldn't check for command-line arguments if the module were imported. (Remember that sys.argv always includes the name of the Python file being executed, so its length is always at least 1.) A Unix-style "usage" message should be printed when an unacceptable number of arguments are provided, and the program should exit. A typical construction is:

```
if __name__ == '__main__':
    if len(sys.argv) < 2:</pre>
        print('Usage: downloadPDFs filename ...')
        for filename in sys.argv[1:]:
            process(filename)
```

sys.modules

This is a dictionary whose keys are the names of the currently loaded modules and whose values are the corresponding module objects. These are the modules that have been loaded into the system. When Python first starts there are 50–75 modules in the system, depending on how it was compiled; the version; and the platform, among other details. Most of the preloaded modules are low-level implementation facilities, but some of them are modules we'll be discussing and you'll be importing. Although these modules are already in Python, their names are not in the interpreter's namespace until you import them. Modules imported by your code get added to this list.

This value is mentioned only as a view into a bit of Python's implementation. You won't do anything with this value, but you can look at it if you're curious. Most importantly, it shows how simple and consistent Python's implementation is: even its collection of modules is an ordinary dictionary. In fact, module objects are themselves very little more than dictionaries.

sys.builtin module names

This is another value that you might find interesting but will rarely need to look at, and certainly will never need to change. It's simply a list of the names of the modules from which the system was built. Most, but not all, are names of modules that are also imported and are therefore in the sys.modules dictionary.

sys.path

This is a list of directories where Python looks for modules when it executes an import statement. By customizing this value, you can inform Python about directories containing Python modules you use in your code (these could be libraries you've downloaded, libraries your organization has built, or libraries of code you've developed yourself).

When IDLE is started, a platform-dependent default directory is added to the front of sys.path—for example, on OS X it would be the user's *Documents* directory. When the interactive interpreter is started from the command line, the command line's "current directory" is added to the front of sys.path. When a Python file is executed, either from the command line or from within IDLE, the file's directory is added to the front of sys.path.

sys.stdin, sys.stdout, sys.stderr

Languages whose programs can be run from a command line generally use three default streams for their input and output, called stdin, stdout, and stderr. As their names suggest, the first is for input, the second is for output, and the third is for error messages. The sys module defines names for these three streams.

The input function always reads from sys.stdin. However, the input stream is not necessarily taken from the user's typing—a file or the output of another process could be redirected to your program, which could read it with input. You could also call sys.stdin.readline or use any of file's other input methods.

The significance of having separate streams for output and error messages is that one or the other can be "redirected" to a file or another program, and it's important to be able to separate normal from error output. In particular, you will often redirect standard output to a file but leave error output printing to the command-line environment, where you can see it immediately. You can use either sys.stdout or sys.stderr as the optional file argument to print, or call any of file's output methods on them.

A subtle difference between sys.stdout and sys.stderr is that sys.stdout is buffered while sys.stderr is not. Low-level operating system input and output operations involve significant overhead. Instead of incurring that overhead for every character or line, output to sys.stdout is accumulated and written only in reasonable-sized chunks. Error messages, however, usually need to be seen immediately, so each line printed to sys.stderr is delivered immediately.

sys.exit([ara])

A function definition can exit a program by calling sys.exit, which returns arg (default 0) to the operating system. (The Unix convention is that 0 signifies no error, 1 signals a runtime error, and 2 indicates a command-line error.)

When a function is defined with a parameter for the stream (open file) the function should print to, that parameter appears in three places:

- In the function definition
- In every call to the function
- In every call to print or any other output function within the definition

Some programs contain many functions that write output to the same destination. It becomes very tedious to have to specify that destination in every function definition, function call, and call to print. An alternative approach is to omit the destination parameter from the function definitions, function calls, and print calls, and let all the output default to sys.stdout. To write the output to the desired file, the program can simply rebind sys.stdout to the result of a call to open. (Except for keywords, there are no sacred names in Python—any name can be rebound with an assignment statement.) If the program will need the original value of sys. stdout, it can save the old value, assign the new value, and, in a finally clause of a try statement, rebind it to its original value. This is shown in the following template.

TEMPLATE

Redirecting to stdout in a Program

You can simplify a portion of a program with a large number of function definitions and calls that all write output to the same destination by rebinding sys.stdout. This is a lot easier than including that output destination as a parameter to every function, an argument in every function call, and a file keyword argument in every call to print.

```
stdout_save = sys.stdout
    sys.stdout = open(filename, 'w')
    ... statements ...
finally:
    sys.stdout = stdout save
```

The system clock: time

The time module does not define any classes. It provides functions for accessing the system's clock, converting among time formats, and working with details of times such as time zones and daylight savings time settings. Most of them are very technical, so we won't discuss them here. A few of the functions, though, are important.



Computer time is defined relative to a zero starting point sometimes called the *epoch*. Different operating systems define the starting point differently. The absolute value is almost never relevant—what is usually significant is the difference between a clock value saved at one point in the program and the clock value sometime later in its execution.

time functions you are likely to find useful include:

time.gmtime(0)

Returns a data structure that describes time zero (in case you are curious)

time.ctime([seconds])

Returns a string showing the date and time at *seconds* past time zero, defaulting to nowtime.time

time.clock()

Returns the number of seconds past time zero, expressed as a floating-point number—use round to convert this value to an int

time.sleep(seconds)

Stops executing for *seconds* amount of time, then resumes

The clock function is useful for timing the execution of Python programs when you are trying to figure out how to make them run faster. The sleep function is used mostly when the program must wait for some external condition. The next template shows how that works.

TEMPLATE

Waiting on an External Condition

Many programs contain a function that must wait until some external condition has been satisfied. A typical example is a file becoming available for reading.

```
while not external-condition-test:
    optional-waiting-action
    time.sleep(seconds)
```

The optional waiting action in the template might be something like printing a period (.) to the terminal to show the user that the program is waiting. Without the delay built into the loop, the program would repeat the test and action immediately over and over again, wasting processing resources that the computer could better use doing something else.

Command-Line Utilities

Next, we'll look at utilities for processing information provided by the command-line invocation of a program. We'll also see some utilities that go in the other direction, invoking command-line commands.

Reading multiple files: fileinput

Many programs you write will take a list of files as their command-line arguments and do the same thing with each of them. While you could write a loop that opens and processes each one, sometimes you'll want to treat the contents of the files as one continuous stream of lines. You may also want information about how many lines have been encountered, which file is being read, and so on. The fileinput module is a convenient tool to use in these situations in place of writing your own code.

The simplest use is:

```
import fileinput
for line in fileinput.input():
    process(line)
```

By default, fileinput.input uses sys.argv[1:] as the list of files to iterate over. However, you can give it a different list as an argument. An unusual feature of this module is that it creates a global state that can be accessed through its functions—you don't even need to create and manage an instance of a class (it does that for you). The state functions include:

```
fileinput.filename()
    Returns the name of the current file
fileinput.filelineno()
    Returns the number of the line just read from of the current file
fileinput.isfirstline()
    Indicates whether the line just read is the first of its file
fileinput.lineno()
    Returns the cumulative line number of the line that has just been read
```

```
fileinput.nextfile()
```

Closes the current file, skipping the rest of its lines

Command-line options: optparse

You will sometimes want your Python code to handle the kinds of command-line options you've seen in programs you've used. Dealing with various option formats and analyzing all of a program's command-line arguments to see which are options, which options they are, and what values (if any) they have can turn into a mind-boggling exercise. Even if you write some simple code to handle a couple of options, you may well add more options in the future, and the code will grow in complexity.

The optparse module does all the command-line argument parsing for you. You just have to tell it what options the program can take and a bit of information about each. When it's done its work, you are left with an object containing values for the options and a separate list of the arguments that follow the options. Also, optparse can generate -h and --help options automatically from the options you describe.

The optparse module implements a large set of features. Just a few of the most important are described here.

TEMPLATE

Using optparse

The basic use of the module follows the outline:

```
import optparse
optparser = optparse.OptionParser(usage=usage string)
optparser.add option(details)
optparser.add option(details)
# ...more options...
optparser.set defaults(option default values as keyword arguments)
options, args = optparser.parse args()
```

The first value returned by parse args is a dictionary-like object whose values can be accessed using keyword notation (i.e., options.key). The other value is a list of the positional arguments that followed all the options on the command line.

The optional usage string is normally in the typical Unix format. For example, a program to copy something from a source to a destination might have a usage string such as the following:

```
'usage: copy [options] source destination'
```

If the program can take an indefinite number of arguments, you would end the string with an ellipsis:

```
'usage: copy [options] filename...'
```

When the program prints the usage string, "prog will be replaced with the name of the program from the command line (sys.argv[0]). (Sometimes it's useful to use filesystem links so you can call the same program with different names and do something different depending on which name is used; that's a reason for using %prog other than just for convenience.) If you don't provide a usage string, a default value will be used:

```
"usage: %prog [options]"
```

The details part of each call to add option provides two kinds of information:

 One or more strings indicating the way the option can be specified on the command line; for example:

```
parser.add option('-o', '--output', '--output filename')
```

• Optional keyword arguments ("attributes") giving further details about the option

Typically, two strings are provided, following the usual Unix style: a "short" name beginning with a single dash, and a "long" name beginning with two dashes. The value of the option specified on the command line (if any) is called the option argument. The arguments of both short and long option names may be separated from the names by whitespace. Arguments of short option names may also appear directly following the option name with no intervening whitespace. A long option name's arguments may be separated from the name by an equals sign, without whitespace.

No other syntax for option arguments is supported. Therefore, the possible ways to provide an option argument on the command line are:

- -o value
- -ovalue
- --option value
- --option=value

The basic keyword arguments are shown in Table 6-3, along with their default values.

Table 6-3. Option parser attributes

Name	Default	Description
action	'store'	What the option does (see below)
type	'string'	Type of argument expected (see below)
nargs	1	For n , the value of the attribute; if $n > 1$, the option takes the next n values from the command line and returns them in a tuple
default	None	The default value for the option
choices		A list of strings that are the valid values for the option
dest	Name of option	The name to use for the option in the option object returned by parse_args
help		Help text for the option

An option's help text should be a string with no line feeds. When help is printed, the module formats the help strings appropriately.

The option's destination is the key that will be used in storing the option's value in the option dictionary. If an option's specification does not include dest, its destination is determined by its names. If there are one or more "long option" names—those beginning with two dashes—the first is used as the destination. If there is no such name, the first "short option" name—which begins with one dash—is used. Default destinations do not include the initial dashes of the option name.

Option default values may be set in one of two ways: you can include default values in the option specifications, or you can call the parser's set defaults method. That method takes a series of keyword arguments. Each keyword is the destination name of an option, and its value is the default to use if the user of the program does not specify a value for that option on the command line. The method must be called before the option specifications are parsed.

Table 6-4 summarizes how each option type is parsed.

Table 6-4. Option parser types

Types	How parsed
string	As is
int	Ox for hexadecimal, int if no prefix
float	float()
choice	Like string; check by calling check_choice()

Table 6-5 describes option parser actions.

Table 6-5. Option parser actions

Action	Consequence
store	The option must be followed by an argument on the command line; the argument will be converted according to the option's type and stored in the option object under the name give by its dest
store_true	No command-line value; store the value True
store_false	No command-line value; store the value False
append	Like store, but the command line can contain multiple occurrences of the option and the dest value will be a tuple

The value of the -h/--help option is generated automatically. The help message will show the usage string (if provided) and the name of each option followed by the value of its help attribute. When a help option is specified on the command line, the program exits after showing the help text. Your program can also call the print help function on the instance of OptionParser, which prints the help text but does not exit the program. If %default appears in an option's help string, the value of the option's default will appear in its place when the help message is printed. Examples 9-4 and 9-5 show the use of optparse in a program to download the links contained in a web page (see "Downloading a Web Page's Linked Files" on page 334).

Command-line commands: subprocess

Sometimes you need a program to execute a command as if it had been typed at the command line. This can be a very powerful way of driving other programs from a Python script, controlling the sequence in which they are called, and coordinating their results. It can also be used to invoke command-line facilities rather than programming their equivalents when they don't exist in Python.

The subprocess module provides sophisticated, highly generalized facilities that can be used in various ways. The following two high-level functions provide access to most of the module's capabilities (in both, command can be either a string or a sequence of strings):

subprocess.getoutput(command)

Executes *command* in a shell and returns its output

```
subprocess.call(command, ...keyword args...)
```

Executes command, waits until it returns, then returns the "success value" of the command



The success value is platform-dependent, but it's usually 0. To fit this into the usual Python-style tests, you should think of the result as indicating whether an error has occurred: 0 means no/false (no error has occurred).

Many optional arguments are available for subprocess.call, but we'll consider only the two most important. Example 6-1, presented momentarily, shows the use of this function to approximate the behavior of the Unix 1s command. A more extensive example follows that one.

Probably the most important keyword argument to consider controls whether *command* is invoked directly—as, for example, when it is simply the name of a program to run—or is executed by a command shell. The difference is important both for the form of *command* and for how it is interpreted when executed. The default is to not execute in a shell; specify the keyword argument shell=True to change that. Here are the key points to keep in mind:

Interpretation of the command argument

If executing directly, *command* should be either a string containing just the program name or a sequence whose first element is the program name (the rest being the arguments for that program).

If executing in a shell, *command* can be specified either as a string or as a sequence. A string will be passed directly to the shell. With a sequence, the first element names the command to execute and the rest of the elements are the arguments to that command.

Contents of the command argument

If executing in a shell, *command* can be anything you can type in a shell. It can include wildcard filenames, environment variables (whose names are preceded by a dollar sign), actions that are commands to the shell itself rather than programs to run, multiple commands separated by semicolons, and so on.

If executing directly, no variables will be expanded or wildcard filenames matched.

The other important keyword argument is cwd, the value of which is a string representing a path to a directory. If a cwd argument is provided, the command (whether in a shell or not) will execute in that directory. Many other keyword arguments are also available, but they are for much more specialized technical uses.

Now, here's the promised approximation of the 1s command (Example 6-1).

Example 6-1. An approximation of the Unix ls command

```
def ls(path ='.', args = ''):
    """Invoke the shell ls command with args on path"""
   subprocess.call('ls' + ' ' + args + ' ' + path), shell=True)
```

This definition accepts only one path, but it could easily be changed to accept either one path or a sequence of paths. The important aspect of this definition is the shell=True argument to subprocess.call. Without that there would be no expansion of wildcards, environment variables, and, in Unix, tildes.

Sometimes it's much easier to use a preexisting command-line command than it would be to write your own program to perform a similar function. Unix provides a huge set of programs to manipulate files and their contents. These are particularly useful when dealing with very large files where reading all the data into Python would be impractical or when you need the efficiency and reliability of built-in programs rather than a Python solution you've written yourself. Also, except for the purposes of experimentation and learning, it's almost never a good idea to code functionality you could find somewhere else—keep in mind that this applies to command-line tools.

Example 6-2 uses the Unix sort command to sort the data from a Rebase file by the second value on each line. (In the "bionet" format that value is the sequence recognized by the enzyme.)

Example 6-2. Using the Unix sort command to sort a file

```
"""Produce a file of enzyme names and cut sites from the Rebase bionet file with
prototypes omitted and 's removed, sorted by recognition site sequences" ""
def sort bionet file(filename = 'data/Rebase/link bionet.txt'):
    cleanfilename = write clean(filename)
    sortedfilename = cleanfilename[:cleanfilename.rfind('.')] + '.sorted'
    # Construct Unix sort command
    command = ('sort -f -k 2 -b -o ' + sortedfilename + ' ' + cleanfilename)
    subprocess.call(command) # tell Unix to execute the command
```

```
def write clean(infilename):
    """Write a copy of file named infilename with prototypes and 's omitted"""
    extpos = infilename.rfind('.')
    outfilename = ((infilename if extpos < 0 else infilename[:extpos]) +</pre>
                     .clean')
    with open(infilename) as infile, open(outfilename, 'w') as outfile:
            write clean lines(infile, outfile)
    return outfilename
def clean line(line):
    """Return line without its prototype (if any) and ^ (if any)"""
    line = line.replace('^', '')
    lpos = line.find('(')
    if lpos >= 0:
        rpos = line.find(')')
        return line[:lpos] + ' '*(rpos - lpos + 1) + line[rpos+1:]
    else:
        return line
def write clean lines(infile, outfile):
    for n in range(11):
        line = infile.readline()
                                                    # skip first 10 lines (introduction)
    while len(line) > 1:
                                                    # stop at empty line or end of file
        outfile.write(clean line(line))
        line = infile.readline()
```

The arguments to the sort command the example constructs have the following meanings:

```
-f
    Ignore case
-k 2
    Sort by the second "field"
```

Ignore leading blanks in the field; the definition of a "field" (unless a separator character is specified with -t) is the transition from nonwhitespace to whitespace, so without b, the lines will be sorted by groups according to increasing length of the enzyme name

-0 The name of the output file

Communications

-b

When an application is used seriously it becomes important to keep track of its status and activity. Python provides a range of modules that can be used for this purpose. These include a very elaborate email module, which we will not talk about; a much simpler way to send email, which we will talk about; and a feature-rich logging facility.

Sending email: smtplib

The straightforward way to send an email message is to use the Simple Mail Transport Protocol (SMTP), which dates from the early 1980s, as implemented by Python's smtp module. This protocol is still used by many email programs. The steps involved are creating an instance of smtplib.SMTP, creating a connection to an email server, providing a username and password if necessary, then sending a message that begins with text in a specific format:

```
To: username, username, ...
From: sender
Subject: optional subject line
Content-Type: text/plain; charset="us-ascii"
Body of message
```

Although the sender and receiver(s) are included in the message text, they must be submitted separately when the message is sent. For most mail servers, especially ones external to your organization, a username and password are required. If your organization's email server can accept SMTP requests without a username and password, they aren't necessary; otherwise, you can set up an account and password for your program to use.

Example 6-3 shows how to put all of this together, including catching the exceptions that typically occur when something goes wrong. While it could be used for many other purposes, the reason we include it here is that it is a good way for a program to email someone when something needs attention, or even just to say that everything's OK.

```
Example 6-3. Sending email
import smtplib
import socket
                                 # just for socket.err
def sendmsg(fromaddr, toaddr,
            username=None, password=None,
            subject='', msg='',
            hostname='localhost', port=25):
   # Destination can be a single username or a sequence of them
   dest = toaddr if type(toaddr) == str else ', '.join(toaddr)
   msg = '''To: {1}
From: {0}
Subject: {2}
Content-Type: text/plain; charset="us-ascii"
'''.format(fromaddr, dest, subject) + msg
   connection = None
        connection = smtplib.SMTP()
        connection.connect(hostname, port)
        if username:
```

```
connection.login(username, password)
    connection.sendmail(fromaddr, dest, msg)
    print('\nMessage sent from {}:\nto: {}'.
          format(fromaddr, dest),
          file=svs.stderr)
except smtplib.SMTPConnectError as err:
    print('Attempted conneection to {} on port {} failed',
          format(hostname, port),
          file=sys.stderr)
except smtplib.SMTPAuthenticationError as err:
    print('Authentication for user', username, 'failed',
          'Invalid username-password combination?',
          sep='\n',
          file=sys.stderr)
except socket.error as err:
    print('Socket error:', err, file=sys.stderr)
finally:
    if connection:
        try:
            connection.quit()
        except smtplib.SMTPServerDisconnected:
                            # can't quit an unconnected SMTP
```

Logging: logging

The logging module provides a wide range of options and facilities. The primary advantage of using a logging facility rather than just writing to a file is that different programs can log to the same file without interfering with each other. Other logging features allow various kinds of control over what is logged and where the output goes.

A fundamental concept of logging is that of a priority level. The logging module uses the levels debug, info, warning, error, and critical. They are defined both as functions and, with all uppercase names, values (e.g., logging.WARNING). Example 6-4 shows a minimal use of the module. We won't go into its many other options and capabilities here, but you should keep the many possibilities in mind when the time comes that you need a logging facility.

```
Example 6-4. A minimal logging example
import datetime
import logging
LOG FILENAME = 'temp/logging.out'
# This directs the logging facility to use the named file and to
# ignore any requests for log entries with a level less than DEBUG
logging.basicConfig(filename=LOG FILENAME,level=logging.INFO)
# This appends an entry to the logfile
logging.info("This message was logged at {}."
              .format(datetime.datetime.now().strftime('%H:%M')))
```

If you examine the contents of the logfile you will notice that each line begins with the log level and the word "root". In more sophisticated uses of the logging facilities, you can set up hierarchies of logs; the name of the log that actually handles the request will then appear in place of "root".

The Filesystem

In this section we'll examine important modules for working with the filesystem.

Operating System Interface: os

The os module contains a few environmental values and operations similar to those in sys. Mostly, though, it provides operations for manipulating files and directories. It implements many low-level operations that you are unlikely to use, but it also contains some that you will use frequently.

The functions in os raise an OSError when given arguments of the wrong type, paths that aren't valid (e.g., a file that doesn't exist), or paths to files or directories that your program doesn't have the necessary privileges to read.

Environment access

Before moving on to the main content of the os module, it is worth mentioning two variables it provides:

os.sep

The string used to separate path components—'/' on Unix-based systems, '\\' on Windows*

os.environ

A dictionary containing the names and values of "environment variables" obtained from the operating system environment from which you started Python; assigning os.environ[varname] will change the value of the environment variable varname outside, as well as inside, Python

You can obtain the value of an environment variable either by accessing os.environ as a dictionary or by using the following function:

os.getenv(varname[, defaultvalue])

A function that returns the value of the environment variable *varname*— i.e., the value of the key varname in the dictionary os.environ; if that key is not in the dictionary the function returns *defaultvalue* if specified, and otherwise None

^{*} Remember that backslashes in strings signify special characters, so including a backslash in a string requires two of them; '\\' is a "special character" like '\t' and '\n'.

Managing files and directories

os provides various functions that mimic the kinds of file and directory management commands available from the operating system command line. The module contains many other functions like these for more technical purposes, but the following sections outline the ones you're most likely to use.

Directories. Common directory-related functions include:

os.getcwd()

Returns the current "working directory" (like the command-line pwd)

os.chdir(path)

Makes *path* the working directory

os.mkdir(path)

Creates a directory at path

os.mkdirs(path)

Creates all the directories along path

os.rmdir(path)

Removes the directory at *path*

os.removedirs(path)

Removes all the directories along path

Files. Useful file-manipulation functions include the following:

os.remove(path)

Removes the file specified by path

os.rename(sourcepath, destpath)

Renames the file at sourcepath to destpath

os.startfile(path)

Opens the file at *path* using its associated application

Directory contents. Two very useful functions in **os** deal with the contents of directories:

os.listdir(path)

This function simply returns a list of the names of the files and directories in the directory specified by its argument. The argument is not optional, though it would be nice if it defaulted to the current directory—i.e., '.'. (That said, it's easy enough to define your own version of listdir that calls os.listdir('.').) Note that the list is ordered arbitrarily; you will often want to sort it.

os.walk(path)

This is almost a facility on its own. Its purpose is to produce the names of all the files and directories at path and below. For each directory encountered starting at path, it produces a tuple of the form:

(directory-path, subdirectory-names, filenames)

The *directory-path* in the tuple is a string indicating the path to the directory starting at the initial path argument. The two lists contain directory names and filenames only, not full paths; if you need paths, you simply add the directorypath to the directory name or filename. The function has several optional parameters, but we'll just deal with the basics here.

The value returned by os.walk is another kind of iterable. This special kind of object is integrated with os.walk in an interesting way: each time around an iteration over the result of os.walk you can remove elements from the subdirectory-names list, and the walk will skip those as it descends into the directory hierarchy. This is useful for ignoring directories such as those beginning with a period or those containing some kind of configuration information that you don't want included in the walk. (The special paths '.' and '..'—current directory and parent directory, respectively—are never included in the walk.)

Example 6-5 demonstrates a simple of use of os.walk. It prints the names of all the directories and filenames in an initial path and below, ignoring any directories whose names begin with a period. The function show in path obtains the information to be printed and show directory contents prints it. Indentation is added by show in path to indicate the "depth" of each file and directory name. (Depth—the number of parent directories up to the original path—is easily calculated by simply counting the number of separator characters in the path.) Using indentation to indicate depth is a common nongraphical technique for showing tree-structured information, such as classical biological taxonomies and filesystem folders. Filesystems are inherently tree-structured because each directory can contain subdirectories.

```
Example 6-5. Showing a file tree
```

```
def show directory contents(dirpath, filenames, level):
   print('
             '*level, dirpath, sep='')
   for name in filenames:
                  '*(level+1), name, sep='')
def show in path(startpath, ignoredots=True):
   print(startpath)
    for path, dirnames, filenames in os.walk(startpath):
        for dirname in dirnames:
           if dirname[0] == '.':
                dirnames.remove(dirname)
        show directory contents(path[len(startpath)+1:],
                                                               # strip dirpath
                                filenames.
                                path.count(os.sep))
```

You might recognize this as a Filtered Do iteration (see "Filter" on page 122 in Chapter 4). It's actually a filtered recursive do, but os.walk is handling the recursion. We could instead write this example using the template shown in the previous chapter, where recursive iterations were discussed, but this is only a simple example; for more complex tasks you'll find os.walk more convenient than writing equivalent code yourself.

Temporary files: tempfile

Creating a new file for output with a name that doesn't conflict with existing names is something programs sometimes do. You could write your own code, generating a numbered filename, checking whether one with that name exists, and incrementing the number until you find a name for which no file exists. However, there are many problems with that naive approach. The module tempfile provides a solid facility that takes care of all the intricacies of creating temporary files. It provides the following functions, among some others that are useful for more technical applications:

tempfile.makestemp([suffix, [prefix, [dir, [text]]]])

Creates and returns the absolute path of a temporary file with a name that includes suffix at the end (default '') and prefix at the beginning (default 'tmp'), located in the specified directory (default as described for tempfile.gettempdir below), and in binary mode unless text is given as True. (The "s" in the name stands for "secure.")

tempfile.makedtemp([suffix, [prefix, [dir]]])

Creates and returns the absolute path of a temporary directory; suffix, prefix, and dir have the same meaning as they do in tempfile.makestemp.

tempfile.gettempdir()

Returns the path of the current default directory for creating temporary files and directories. This may be set by assigning tempfile.tempdir. If that is unbound or None, the environment variables TMPDIR, TEMP, and TMP are examined, with the value of the first one that has a value being used. If none of those environment variables has a value, on Windows the function will look for the directories C:\TEMP, C:\TMP, \TEMP, and \TMP; on other platforms it will look for /tmp, /var/tmp, and /usr/tmp.



While these are called "temporary files," that is more a reflection of their intended use than of any special treatment they receive. It is up to the program(mer) to delete temporary files and directories that are no longer needed.

Manipulating Paths: os.path

The os module contains a submodule, os.path, which contains functions for manipulating paths (as opposed to the files and directories at those paths). The following sections outline some of the more important of those functions.

Path components

One of the most frequent kinds of manipulations you will do to file paths is breaking them apart into their components in various ways. The os module provides these functions for that purpose:

os.path.split(path)

Returns a pair (head, tail), where tail is the file part of the path, if any, and head is the directory part of the path, if any (i.e., everything through the last slash is part of *head*, and the rest of *path* is part of *tail*)

os.path.dirname(path)

Returns the file part of the path (the same as the first element of the pair returned by split)

os.path.basename(path)

Returns the directory part of the path (the same as the second element of the pair returned by split)

os.path.splitdrive(path)

Returns a pair (drive, tail), where drive is the drive specification, if any, and tail is the remainder of the path

os.path.splitext(path1)

Returns a pair (path2, ext); if path1 contains one or more periods (.), path2 will be the part of path1 up to the last period and ext the part after the last period; otherwise, the result is (path1, '')

As we saw, the function os.listdir(dirpath) returns a list of the names of all the files and directories that are in the directory at dirpath. The result is often hard to read, partly because it's a list, not the formatted output printed by command-line utilities, and partly because it contains a lot of uninteresting files, such as editor backup files, .pvc files, and so on.

The first problem can be solved simply by using print on each element of the list returned by os.listdir. The second problem can be solved by using os.path.splitext to filter out files with extensions you're not interested in seeing. These solutions are incorporated into the function defined in Example 6-6, a Filtered Do (see "Filter" on page 122 in Chapter 4).

Example 6-6. Filtered directory listing

```
def filtered directory listing(dirpath = '.',
                               ignore extensions = ('.pyc', '.bak')):
   for filename in os.listdir(dirpath):
        if os.path.splitext(filename)[1] not in ignore extensions:
            print(filename)
```

Path manipulations

Many programs deal with paths that have no or only a few directory names, but need full paths for certain purposes. The following functions expand or join paths in ways that would be difficult, or at least laborious, to program yourself:

os.path.abspath(path)

Returns a full absolute version of path

```
os.path.expanduser(path)
```

Returns *path* with an initial ~ or ~*username* replaced by the user's home directory

```
os.path.join(path1, path2, ...)
```

Returns a path formed by joining the arguments appropriately; roughly similar to the expression os.sep.join(path1, path2, ...) (os.sep is the platform-specific path separator mentioned a few pages back) except it doesn't add separators after arguments that already end in a separator, and any argument that is an absolute path causes it to ignore the arguments that precede it

The functions in the os.path module are important tools. Many programs perform at least a few manipulations on file and directory paths. In addition, you can use the os.path functions to define some utilities for your own convenience when using the interpreter. Here are a few examples:

```
def cd(path):
    """Make path the current directory, expanding environment
    variables and a ~ representing the user's home directory"""
    os.chdir(os.path.expandvars(os.path.expanduser(path)))

def merge_ext(path, ext=''):
    """Return path with its extension replaced by ext, which normally
    starts with a period; with no arguments, remove the extension"""
    splitpath, splitname = os.path.split(path)
    return os.path.join(splitpath, os.path.splitext(splitname)[0] + ext)
```

Path information

Often, you just need to test something about a path before doing something with it. Here are three predicates and a function to get the size of a file:

```
os.path.exists(path)
Returns True if there is there a file or directory at path
os.path.isfile(path)
Returns True if there is there a file at path
os.path.isdir(path)
```

Returns **True** if there is there a directory at *path*

```
os.path.getsize(path)
```

Returns the size in bytes of the file at *path*

Example 6-7 shows an example of a Filtered Combine iteration (see "Filter" on page 122 in Chapter 4) in a complete program. The collection is the list of file and directory names in a path; the test is os.path.isfile; and what is being combined is the result of a call to os.path.getsize. To emphasize how brief well-written code can be, most of the examples in this book omit the imports and the "if main" code at the end. They are included in this example, though, so you can see what it's like to turn a simple function like this into a usable program.

Example 6-7. Calculating the total size of files in a directory

""Print the sum of the sizes of the files in the directory given on the command line (or the current directory if none is given), in megabytes rounded to 2 places"""

```
import os.path
def directory size(path='.'):
    """Sum of the sizes of all files in the directory at path, including
    those beginning with a '.', and ignoring subdirectories"""
                                                      # identity element
    for item in os.listdir(path):
        if os.path.isfile(item):
                                                      # the test
            result += os.path.getsize(os.path.join(path, item))
    return result
if name == ' main ':
    if len(sys.argv) > 2:
        print('Usage: directory size path')
        size = directory size('.' if len(sys.argv) == 1 else sys.argv[1])
        print(round(size/1024/1024, 2), 'Mb', sep='')
```

Another very useful utility would be one that prints a directory structure in hierarchical form. We can define one based on the Recursive Tree Iteration template of Chapter 4 (see "Recursive iterations" on page 128). However, whereas the template assumed the first argument was an entire tree, in this example each call to the function provides just a path. The function gets the tree below that path from a call to os.listdir. Example 6-8 shows the code.

```
Example 6-8. Hierarchical directory listing
```

```
def dirtree(path='.', ignoredots=True, level=0):
   print path(path, level)
                                                      # "do something" with tree root and level
   for name in os.listdir(path):
                                                      # repeat with the rest of the tree
        subpath = os.path.join(path, name)
        if os.path.isdir(subpath):
            dirtree(subpath, ignoredots, level+1)
def print path(path, level):
   print(' ' * 3 * level, path, sep='')
```

Filename Expansion: fnmatch and glob

The fnmatch and glob modules reproduce the wildcard expansion operations of command-line shells. The difference is that fnmatch works directly on strings, while glob actually goes out to the indicated path to expand wildcards relative to the filesystem content. Both use the command-line matching syntax shown in Table 6-6.

Table 6-6. Wildcard characters for fnmatch and glob

Pattern	Meaning
*	Match 0 or more characters
?	Match a single character
[characters]	$Match \ any \ of the \ characters \ inside \ the \ brackets; the \ characters \ are \ not \ separated \ (e.g., \ '\ [aeiou]')$
[!characters]	Match any of the characters except those inside the brackets

fnmatch

The main functions of the fnmatch module are as follows:

```
fnmatch.fnmatch(filename, pattern)
    Returns True if the string filename matches the string pattern
fnmatch.filter(filenames, pattern)
```

Returns the list of strings in *filenames* that match the string *pattern*



The fnmatch module is extremely simple: it treats periods and slashes the same as any other characters. For example, '.pythonrc' matches '*rc'. This would not be the case on the command line.

Example 6-9 gives an example of using fnmatch.filter in a Collection Combine (see Example 4-15, shown in the section "Combine" on page 117) to search a directory hierarchy for all files whose names match a particular pattern. You could use this to locate all files with a particular extension (such as .fasta), all files with a particular prefix (such as gb), and so on.

Example 6-9. Find files matching a pattern

```
def find matching files(startpath, pattern):
    """Return a list of filenames that match pattern in the directory tree starting at startpath"""
    paths = []
    for path, dirnames, filenames in os.walk(startpath):
        for dirname in dirnames:
            if dirname[0] == '.':
                dirnames.remove(dirname)
        paths += [os.path.join(path, filename)
                       for filename in fnmatch.filter(filenames, pattern)]
    return path
```

alob

More useful is the glob module. It uses the same pattern syntax but matches against the contents of the filesystem itself, not against a list of strings the way the fnmatch functions do. glob matches the way the command line does: it ignores files beginning with a period (unless the pattern begins with a period), and it treats slashes as separating path components.



The glob functions do not expand tildes and environment variable names in paths; for that you need os.path.expandvars.

The glob module can return results either as a list or as a generator, depending on which function you call. The generator version is important for when there are a very large number of filenames matching the pattern, only some of which need to be used. The two relevant functions are:

glob.glob(pattern)

Returns a list of paths (strings) that match *pattern* (which doesn't necessarily contain wildcards)

glob.iglob(pattern)

Same as glob.glob, but returns an iterator instead of a list

For example, to do something to each Python file in the current directory:

```
for filename in glob.iglob('*.py'):
```

Shell Utilities: shutil

The shutil module provides functions for performing common file operations at a higher level than those provided by os—copying and deleting, in particular. There are several kinds of copy operations in the module, but the ones you would normally use are as follows:

copy(source, destination)

Copies the file specified by the path *source* to the file or directory specified by the path destination; if destination is a directory, the file will be copied into it



Like its command-line equivalent, copy will overwrite a file of the same name as the one specified by source—either a file by that name in destination, if it's a directory, or in the directory containing destination, if it's a file.

copytree(source, destination)

Recursively copies the directory *source* to *destination*; all parent directories in the path destination will be created as necessary and an OSError is raised if destination already exists

Moving and removing operations are:

move(source, destination)

Moves the file or directory *source* to *destination*

rmtree(path)

Deletes the directory *path* and everything under it



When programming with these functions you should test your code on directories and files set up for that purpose, so you don't end up unintentionally copying or deleting files.

Comparing Files and Directories

You may be familiar with using a feature of an editor or even a separate application to compare the contents of two files or two directories. Operations like these can have many practical uses in programs, too. For example, you may need to extract the differences between two versions of a program's hourly or daily output, or you might need to know if the results of some BLAST queries you've run are any different from the previous results. Directory comparison is similarly useful: you can compare the contents of directories that contain files produced by programs you are using and extract or report the differences.

File and directory comparison: filecmp

The filecmp module provides just two functions, along with a class having only a few methods, but they are powerful and easy to use:

filecmp.cmp(filepath1, filepath2)

Compares the files at filepath1 and filename2 and returns True if their contents are equal

filecmp.cmpfiles(directorypath1, directorypath2, filepaths)

Compares the files in the list *filepaths* in the directory at *directorypath1* with the corresponding files in the directory at *directorypath2*, and returns three lists of file paths:

matches

The paths in *filepaths* to files whose contents were the same in both directories

mismatches

The paths in *filepaths* to files that were in both directories but had different contents

errors

The paths to files in *filepaths* that were not found in both directories or that caused some kind of error when an attempt was made to read them (e.g., because of inadequate user permissions)

filecmp.dircmp(directorypath1, directorypath2, hidenames)

Creates an instance of the class filecmp.dircmp to compare the directories at directorypath1 and directorypath2, with hidenames a list of names to ignore (defaulting to [os.curdir, os.pardir])

Instances of filecmp.dircmp implement the following methods that print fairly elaborate reports to sys.stdout:

report()

Prints a comparison between the two directories

report partial closure()

Prints a comparison of the two directories as well as of the immediate subdirectories of the two directories

report full closure()

Prints a comparison of the two directories, all of their subdirectories, all the subdirectories of those subdirectories, and so on (i.e., recursively)

In addition, many details of the comparisons that the reporting methods print out may be accessed directly as attributes. Their names use the syllable "left" for what was found in directorypath1 and "right" for what was found in directorypath2. The attributes are:

left list

The names of files and subdirectories found in directorypath1, not including elements of *hidelist*

right list

The names of files and subdirectories found in directorypath2, not including elements of hidelist

common

The names of files and subdirectories that are in both directorypath1 and directorypath2

left only

The names of files and subdirectories that are in *directorypath1* only

right only

The names of files and subdirectories that are in *directorypath2* only

The names of subdirectories that are in both *directorypath1* and *directorypath2* common files

The names of files that are in both directorypath1 and directorypath2

common funny

Names common to both *directorypath1* and *directorypath2* but that name a file in one and a directory in the other, along with names of files and directories that caused an error when an attempt to read them was made

same files

The paths to files whose contents are identical in both *directorypath1* and *directorypath2*

diff files

The paths to files that are in both *directorypath1* and *directorypath2* but whose contents differ

funny files

The paths to files that are in both *directorypath1* and *directorypath2* but could not be compared for some reason

subdirs

A dictionary that maps names in common_dirs to dircmp objects

Showing the differences between two files: difflib

Suppose you are faced with two large FASTA files that you know differ only slightly. You may know how to use your editor, the Unix diff command, or another application to get details about the differences between the two files. But what if getting those differences is part of a series of steps your program must perform that at some point involves the difference between the two files. You could use subprocess.getoutput, giving as its argument a command you construct, but then you would have to write code to parse the result it returns. What you really should use is Python's difflib module. It provides classes for three different kinds of comparisons:

difflib.SequenceMatcher

Finds the differences between two hashable sequences (i.e., strings, tuples, tuples of strings, etc.). It provides the following instance methods:

```
set_seqs(seq1, seq2)
set_seq1(seq1)
set seq2(seq2)
```

Set the sequences to be compared.

find longest match(start1, end1, start2, end2)

Finds the longest sequence common to <code>seq1[start1:end1]</code> and <code>seq2[start2:end2]</code>, returning three values: the position in <code>seq1</code> where the common sequence began, the position in <code>seq2</code> where the common sequence began, and the length of the common sequence

get_matching_blocks()

Returns a list of triples describing all the matching subsequences of *seq1* and *seq2* in the form returned by find longest match

difflib.Differ

Compares two sequences of lines of text, finding both completely different lines and differences within a pair of lines using the following method:

compare(stringlist1, stringlist2)

Compares the sequences of strings stringlist1 and stringlist2 and returns a list of lines showing which are the same in both lists, which appear only in stringlist1, which appear only in stringlist2, and which appear in both but with slight differences (which are marked in the result returned)

difflib.HtmlDiff

Like difflib.Differ, but prepares HTML-formatted output according to specified parameters. It provides two methods:

```
make file(stringlist1, stringlist2 [, title1=''[,title2='']])
```

Returns a string containing the HTML for a table highlighting both the interline and intraline differences between the strings in stringlist1 and those in stringlist2. (The function's name is a misnomer: it doesn't write to a file, it returns a string.) The optional parameters title1 and title2 are the strings to use as column headers in the HTML table.

```
make table(stringlist1, stringlist2 [, title1=''[,title2='']])
```

Works just like make file, except that it only returns the HTML for the difference table itself, not the surrounding text that would make it a complete HTML page.

There are further functions in difflib and other methods supported by its three classes. These can provide extensive control over exactly what is matched and how it is reported.

Working with Text

Next, we'll look at some modules for text manipulation. Three address specific needs. A fourth supports reading from and writing to strings instead of files, a feature with wide applicability.

Formatting Blocks of Text: textwrap

The textwrap module provides functions for formatting blocks of text. Programming such operations is notoriously tedious and error-prone, so it's good to have these functions available when you need them:

textwrap.dedent(text)

Returns a copy of the string text with whitespace common to all of its lines removed; for instance, if every line begins with four or more spaces, the result will be shifted to the left four spaces. While tabs and spaces are both whitespace, this function does not treat them as matching.

```
textwrap.wrap(text, [width[, ...]])
```

Wraps the string text as a single paragraph, returning a list of lines (without newline characters) of at most width characters (default 78).

```
textwrap.fill(text, [width[, ...]])
```

Wraps the string text as a single paragraph, returning a single string containing the wrapped paragraph (including newlines). This is just shorthand for:

```
'\n'.join(textwrap.wrap(text))
```

In addition to width, both wrap and fill accept a large number of keyword arguments for controlling their behavior:

```
expand tabs (default True)
```

If true, use text.expandtabs in place of text

```
replace whitespace (default True)
```

If true, replace every whitespace character with a space; if expand tabs is false each tab character is replaced with just one space, but if it's true tabs are expanded before whitespace is replaced

drop whitespace (default True)

If true, eliminate whitespace that appears at the beginning and ends of lines after wrapping, except for whitespace at the beginning of the first line

```
initial indent (default '')
```

Specifies a string to add at the beginning of the first line

```
subsequent indent (default '')
```

Specifies a string to add at the beginning of every line except the first

fix sentence endings (default False)

If true, tries to identify sentence endings and ensures that sentences are separated by exactly two spaces

```
break long words (default True)
```

If true, words longer than width will be broken so that no line is longer than width

break on hyphens (default True)

If true, lines may be broken at hyphens in compound words in addition to being broken at whitespace

The textwrap functions fill and wrap are particularly useful for breaking up a sequence into lines of a fixed length, as when writing a FASTA file. Given a sequence string named seq, we can divide it into a list of lines using wrap or keep it as a long string but with newline characters inserted at appropriate positions using fill:

```
textwrap.wrap(seq, 80)
                                   # break up seq to a list of 80-line strings
textwrap.fill(seq, 80)
                                   # insert newlines every 80 characters
```

Of course, like all string manipulations in Python and its library modules, textwrap.wrap returns a new string.

String Utilities: string

An assortment of string-related values, functions, and classes are collected in the string module. This module has nothing to do with the implementation of the builtin str type; it provides strings that describe kinds of characters and a template facility. Useful string module values include:

```
string.whitespace
   Space, tab, line feed, return, form feed, and vertical tab
string.punctuation
   Punctuation characters as defined in the local character set
string.digits
    '0123456789'
string.hexdigits
    '0123456789abcdefABCDEF'
string.ascii lowercase
    'abcdefghijklmnopgrstuvwxyz'
string.ascii uppercase
    'ABCDEFGHIJKLMNOPQRSTUVWXYZ'
string.ascii letters
    string.ascii lowercase + string.ascii uppercase
string.printable
   whitespace + punctuation + digits + ascii letters
The module has one function that will occasionally be useful:
string.capwords(string)
```

Returns a new *string* formed by splitting *string* at whitespace, capitalizing each of the words, and joining them. Its definition is:

```
def capwords(s, sep=None):
    return (sep or ' ').join([x.capitalize() for x in s.split(sep)]
```

The module's template facility is easier to use than str.format with large bodies of mostly constant text (e.g., a generic web page with placeholders for a few details). Substitution is done according to a dictionary. Wherever a placeholder of the form \$key appears in the template, the value of key in the dictionary is substituted. Keys must be valid identifiers (i.e., names you could give to values, functions, etc.). \$\$ is replaced by a single \$. In contexts where \$key is followed immediately by characters that could be part of a key, \${key} is used instead.

A template is created as an instance of the class Template, with the template string given as an argument. To make the substitutions on the template instance's string, the method substitute is called on the instance with a dictionary as its argument. If any of the keys are missing from the dictionary, substitute raises a KeyError exception. The

method safe substitute is similar, but instead of raising an exception for missing keys it leaves the original placeholder intact. There is no provision for deleting placeholders that aren't keys in the dictionary.

Example 6-10 shows a function that demonstrates the use of templates for a simple form-letter facility. It assumes there are a number of files with similar placeholders. The function's parameters are a subject string, the name of the file containing the template, a substitutions dictionary, and a list of email addresses. The function creates a template from the contents of the designated file, makes substitutions using the dictionary, and for each email address calls the function sendmsg, defined in Example 6-3.

Example 6-10. Using the string template facility to email a form letter

```
def broadcast(subject, filename, substitutions, receivers):
   with open(filename) as file:
        message = string.Template(file.read()).substitute(substitutions)
   for receiver in receivers:
        sendmsg('your-program',
                receiver,
                subject=subject,
                msg=message
```

Comma- and Tab-Separated Formats: csv

Comma-separated values (CSV) files are a widely used "interchange" format, especially in conjunction with spreadsheet and data-collection applications. The term actually applies to a variety of text field conventions, including tab-separated. The exact rules for representing a field's value are not standardized and can vary in frustrating ways from one application to another. Python's csv module hides this variability from you, allowing you to focus on the real purpose of your program.

Very briefly, the module contains the functions csv.reader and csv.writer. It also includes facilities for defining other formats and for more complicated reading and writing operations, but we won't cover those details here. Here are the basics:

```
csv.reader(source[, dialect='excel'])
```

Returns an iterator for the lines of *source* (which can be any iterable that produces strings; if a file, it must already be open).



When opening a file to use with csv.reader, it is necessary to provide a keyword argument in the call to open for a parameter that wasn't included in the description of the function in Chapter 3: newline=''. This allows the reader to correctly interpret newline characters inside quoted field values in a platform-independent way.

Each step of the iterator (either one cycle of a for statement or the result returned by a call to next) produces a list of the values in the comma-separated fields of one line.

If dialect is 'excel-tab', reads the file as tab-separated values; otherwise, reads it as comma-separated values.

```
csv.writer(destination[, dialect='excel'])
```

Returns an instance of a csvwriter for a file—or anything else with a write method—that is open for writing.

If dialect is 'excel-tab', writes tab-separated values; otherwise, writes commaseparated values.

The writer provides the following methods (the "row" in their names refers to the widespread use of CSV files as representations of simple spreadsheets, with each list of fields corresponding to a spreadsheet row):

writerow(fieldlist)

Writes the fields of *fieldlist* to the writer's *destination* according to the writer's dialect, followed by '\n'

writerows(rowlist)

Does csywriter.writerow for each row in rowlist

String-Based Reading and Writing: io

The io module provides the core implementation of Python's input and output facilities, including the definition of open. It offers classes for a variety of capabilities. Some of them can be used for both reading and writing, while others are for just one or the other.

StringIO is used for reading from and writing to strings, and ByteIO is for reading from and writing to bytearrays. This allows you to work with data in memory just as you would with a file on the disk, with the same set of methods (such as readline and writeline). You can also provide instances of these classes as the optional file parameter in calls to print.

To write to a string, you create an instance of io. StringIO, generally with no arguments. That object contains an internal buffer, which is the destination for the strings written to it. The method getvalue obtains a string based on the contents of that buffer. You can call it at any time, even multiple times, until close is called for the instance. (You can either call close explicitly in a finally clause of a try statement, or simply use a with statement, which will call close for you the way it does with files.)

TEMPLATE

String Output

```
import io
with io.StringIO() as dest:
                                       # create new StringIO
    write to dest using methods such as write and writelines
    result = dest.getvalue()
                                      # get the string from dest
```

Instances of io. StringIO are also used for reading from strings. The difference is that when you create the instance you provide a string as an argument. Subsequently, the io.StringIO instance can perform the usual input operations, such as readline and readlines.

TEMPLATE

String Input

import io with io.StringIO(string) as src: # string is the StringIO source read from src using methods such as readline and readlines

With io.StringIO, you can use a string as the source in place of a file in code such as the definition of get items shown in various earlier examples. Where a function uses open and passes the open file to get items, it could just as well create an io.StringIO from a string and pass that instance to get items. This would work with most of the examples in Chapters 3 and 4 that used readline to parse the contents of a file.

Persistent Storage

Data stored in text files is adequate for many applications. However, as the quantity and complexity of the data increases, it becomes too cumbersome to rely on formatted text. This is especially true when an application selectively accesses small amounts of data from a much larger store. For these cases, Python provides three persistent storage modules that use special binary files to store objects more like the way they are represented in memory. These mechanisms make access to external data much faster and much easier to program.

Persistent Text: dbm

The simplest form of persistent storage is essentially a file-based dictionary whose keys and values are both text. This is implemented by the dbm module, which provides the following function for opening database files:

dbm.open(filename[, flaq])

Returns an object providing access to the database in filename; flag must be one of the following:

'r'

Opens an existing database for reading (the default)

'w'

Opens an existing database for reading and writing

'c'

Opens a database for reading and writing, creating it if it doesn't exist

'n'

Opens a new, empty database for reading and writing, replacing it if it already exists

Only one program at a time can have a dbm database open (with an exception shown for the gnu variation discussed momentarily). Table 6-7 shows the operations and methods supported for an object returned from dbm.open.

Table 6-7. Core dbm operations and methods

Operation/method	Description
db[str1] = str2	Stores $str2$ as the value of key $str1$ (as bytes)
db[str1]	Returns the value of key str1 (as bytes)
<pre>del db[str1]</pre>	Deletes the entry for the key str1
str1 in db	Returns true if db contains the key str1
str1 not in db	Returns true if db does not contain the key str1
db.keys()	Returns a list of all the keys in db

There are three submodules of dbm, each of which implements the core functionality in a different way. When you create a new file with dbm.open it determines which submodule's open function to use based on what facilities are available in the operating system environment. † When dbm. open is called on an existing file, it determines which of the implementations created it and calls the corresponding submodule's open function. (You can tell by the type of the object returned which module was used; you can also specify which to use by importing a submodule and calling its open function.) The submodules are:

dbm.ndbm

An implementation based on the Unix (n)dbm facility

[†] Python running on Windows will have only dbm.dumb. Otherwise, dbm.ndbm should be present as well. When installed through a Linux package manager, Python may have dbm.gnu. The dbm.gnu implmentation will also be available in Python built from its source files with the gdbm library installed.

dbm.gnu

An implementation based on GNU gdbm

dbm.dumb

A pure Python implementation

Beyond their implementation of the core operations shown in Table 6-7, the submodules differ somewhat. The current implementation of dbm.dumb.open ignores its flag parameter: files are always opened for reading and writing and created if they don't already exist. The ndbm implementation provides two additional methods that are just like the ones with the same names provided by dict: get and setdefault.

The gnu variation is the richest of the three. The flag parameter of gnu.open can have additional characters that specify further details of how the database file will be used:

'f'

Opens an existing database in "fast mode"—individual changes to the database are not immediately written to the file but rather are buffered for efficiency, similar to the way the buffered output stream sys.stdout works

's'

Opens an existing database in "synchronized mode"—each individual change is immediately written to the database file, similar to the way the unbuffered output stream sys.stderr works

'11'

Doesn't lock the database, allowing other programs and other users to access its contents while your program has it open

The gnu submodule also provides some additional features. The method reorganize is used to compact the database file after many items have been deleted. If the database has been opened with 'f' included in the flag argument, the method sync forces accumulated changes to be written out to the file.

The firstkey and nextkey methods allow you to iterate using the keys of a gnu database. Start with a call to firstkey, which returns a key. Then call nextkey with the first key to get the next one. Repeat the process with the result of the previous call to nextkey until nextkey returns None.

The Rebase data reader example at the end of Chapter 4 constructed a dictionary whose keys were the names of restriction enzymes and whose values were recognition sequences. It included functions to write the parsed data to a file in a simple format and to read the data from that file into a dictionary. With a few small changes, we could convert that program to use dbm instead of a dictionary to store its data. This would mean the data would be loaded only once; subsequently, individual entries could be accessed directly from the database file, and programs that use the database wouldn't need the reading and writing functions at all.



The datafile used for this example is fairly small, so in practice switching to dbm wouldn't make much of a difference. However, if the datafile were much larger it could be significantly more efficient to load the data just once and then access it directly from the database. Even if the file is relatively small, it can be advantageous for other programs to use dbm to access the data so that they won't need the original parsing code. (As always, we want a given piece of code confined to one place, especially one as complex as the parsing code from Chapter 4's examples. In fact, it would be a good idea in this case to have the code that loads the database in a separate file from the code that uses it.)

Example 6-11 shows the functions from that example for loading the data into the table, with modifications to use dbm. The functions that parse the original file will be unaffected by the switch to dbm, but they will be used only once, to load the database. Note that dbm files must be closed, and they must be closed explicitly—they don't support the automatic closing feature of with statements.

Example 6-11. Loading Rebase data into a dbm database

```
import dbm
def load enzyme data(data filename, dbm filename):
                                                              # 'n' for new database
        table = dbm.open(dbm filename, 'n')
        with open(data filename) as datafile:
            load enzyme data into table(datafile, table)
    finally:
        table.close()
def load enzyme data into table(datafile, table):
    line = get first line(datafile)
    while not end_of_data(line):
        key, value = parse(line)
        store entry(table, key, value)
        line = get next line(datafile)
    # return table not needed
def store entry(table, key, value):
    table[key] = val
                                             # table is an open dbm file, but [] unchanged
```

After loading the data into the database by running this program, individual entries would be accessed as follows:

```
>>> table = dbm.open(dbm filename, 'r')
>>> table['AatII']
b'GACGT^C'
>>> table.close()
                                           # can do repeated reads before closing
```

Notice that the data returned is a bytes object, not a string. The database stores both keys and values as bytes. Strings are automatically converted into bytes when stored in the database, but not when they are retrieved.



To convert a bytes object to a str you need to call its decode method. If you want to use an encoding other than the default, you should convert your keys and values to bytes explicitly when storing them into the database. To convert a string s to bytes for a chosen value of encoding, (e.g., 'utf-8'), call s.encode(encoding) when storing it into the database. To convert a bytes b to a string using a specific encoding, call b.decode(encoding).

Persistent Objects: pickle

Using disk files to store bytes is straightforward, since bytes are exactly what files contain. All the dbm module adds to the raw bytes is a bit of structuring so it can efficiently access values by their keys, the way dictionaries do. (Some versions of dbm use a separate file for an index into the file.) This simple mechanism is fine if all you need is a persistent dictionary with keys and values stored as bytes. Storing anything more complicated presents a number of problems, though, some of them quite deep.

The Python module pickle[‡] solves some of those problems. It encodes objects into byte representations from which the objects can be re-created. This facility can handle selfreferential data structures, references to types and functions, large objects, and objects with intricate internal structures. The module's functions read and write byte-encoded representations of objects, but you don't need to know anything about those representations—the module takes care of all the details for you.



Pickling is strictly byte-based; therefore, files used to write and read pickled objects must be opened in binary mode. This doesn't affect the programs: unlike dbm, pickle restores objects to their original types. In particular, stored strings are restored with type str, not bytes.

When an object is encoded, any object that is the value of one of its attributes will also be encoded, along with references to classes and functions. Therefore, pickling one object may actually result in the encoding of many objects. It is often convenient to pickle an entire list or dictionary, which in turn pickles all the list elements or dictionary keys and values.

The pickle module's core functions are:

pickle.dumps(object)

Produces a byte-encoded representation of *object* that pickle.loads can use to create an equivalent object

[‡] Other languages, libraries, and books use other terms for what Python calls "pickling," including "marshaling," "serializing," "flattening," or "persisting."

pickle.loads(bytesinstance)

Re-creates the object represented by bytesinstance, such as that returned by pickle.dumps

pickle.dump(object, bytestream)

Pickles an object and writes it to bytestream, which must be a binary write stream such as a file opened with mode 'wb' or an instance of ByteI0; more than one object may be pickled and written to the same byte stream

pickle.load(bytestream)

Reads an object from bytestream, which must be a binary read stream such as a file opened with mode 'rb' or an instance of ByteIO, and unpickles it; bytestream may contain a series of pickled objects, in which case each call to pickle.load reads just one

TEMPLATE

Using pickle

To save an object using pickle:

with open(filename, 'wb') as file: pickle.dump(object, file)

To load an object from a pickle file:

with open(filename, 'rb') as file: result = pickle.load(file)

An important use of pickling, though we won't demonstrate it, is for communicating objects across networks. There is an example in Chapter 9 (see "Server fundamentals" on page 338) that demonstrates sending simple text between a client and a server. Much more complex architectures can involve cooperating processes on multiple machines rather than a simple client/server configuration, in what's known as distributed computing. Typically, these processes must exchange more complex data. If the processes are running Python programs they can send (copies of) groups of objects among each other by pickling them and sending the resulting text, which can then be unpickled by one or more receivers.

Keyed Persistent Object Storage: shelve

One of the limitations of pickling objects to files is that the code loading the objects must know what to do with what it loads. This isn't a problem if the file contains just one object, including the case where what was pickled was a single collection. With many separate objects pickled to a file, it would certainly be possible to retrieve them one at a time in a loop, collecting them into a list or dictionary. However, this would not be a useful approach in situations where only one of many objects is needed.

Pickling a collection of objects also has its own problems. A dictionary with many thousands of list values may be used by a number of small applications. Unpickling the entire collection of objects each time a program starts could lead to long delays, in addition to wasting a great deal of memory space when only a few instances of the classes are actually needed. The same problems reappear if new instances are created or old ones changed: it will be necessary to repickle the whole collection.

The solution to these problems is to pickle each object individually, rather than pickling whole collections of objects, and to identify each one by a name. Instead of pickling and unpickling the entire dictionary, this approach will allow an application to access just the values it needs using their names. As with so many other things in Python, the way to do this is with a dictionary mechanism. The module shelve combines the kevedstorage aspect of dbm with the object persistence capabilities of pickle. Keys must be instances of str (not bytes). The values may be anything that can be pickled. Like dbm, the shelve module provides a function for opening database files:

```
shelve.open(filename[, flag='c'])
```

Opens *filename* as a persistent dictionary (i.e., a dictionary whose keys are strings and whose values are pickled objects); flag has the same options as for dbm.open (see "Persistent Text: dbm" on page 243)

An opened "shelf" acts like a dictionary, with one critical distinction: modifying a value obtained from a shelf does not affect the value stored in the shelf file. The only actions that affect the file are assignments and deletions—i.e., db[key]=value and del db[key]. If you want to modify a value that is in the file, you must explicitly use an assignment statement with the modified value.

TEMPLATE

Using shelve

Assignment of a shelf entry does the pickling for you, so this module is very easy to use.

```
db = shelve.open(filename, 'c')
db['key'] = value
                          # store value in the shelf, naming it key
db['key']
                          # get the value with name key
del db['key']
                          # remove the entry with name key
db.close()
```

Debugging Tools

Debuggers are facilities that allow you to investigate the state of your program when its execution has stopped. You can use them to determine the sequence of function and method calls that led to the current place in the program, look at the values of local names, execute assignment statements to rebind local names, and "step" through your code to observe what happens.

There are two kinds of Python debugging facilities: the pdb module and tools built into IDEs, including IDLE.

The Python debugger: pdb

Whether from the command-line interpreter, IDLE's Python Shell window, or even a Python file, importing the module pdb gives you access to some highly useful debugging tools. The debugger uses its own read-evaluate-print loop, with the prompt (pdb) and some different behavior than is found at Python's usual top level:

- In general, you can type anything to the debugging prompt that you can type to Python, with the same effect.
- In addition, there is set of *commands*—not function calls—that you can type to perform various actions specific to debugging.

The pdb module provides three functions for entering the debugger:

- Immediately after a traceback, before doing anything else, call pdb.pm0 ("postmortem"). That will put you into the pdb command loop, positioned at the location of the error that caused the traceback.
- Put a call to pdb.set trace0 in your code. When Python executes that line, it enters the pdb command loop.
- Call pdb.run(), giving it a string as an argument. The string would be something you would type to the interpreter. The debugger pauses immediately, and then you can step through your code.

You can also run the debugger on a Python file without importing the module by typing the following at the command line:

```
python3 -m pdb file.py
```

This will enter the command loop before the program even starts running.



The debugger is a tool that warrants continual experimentation and learning. As you become more comfortable with what you learn initially, you can explore more powerful techniques and advanced commands.

Following are descriptions of the basic commands of the pdb command loop; you can type just their first letter or the entire word. First we'll look at commands for getting information and quitting:

h(elp) [command]

Without an argument, prints the list of available commands. With an argument, prints help about that command. If the argument is pdb, prints the full documentation.

w(here)

Prints a full "stack trace," with the most recent frame at the bottom and an arrow showing the current context for other commands. This is like what you see in a traceback, but it shows the entire stack, not just the last few entries.

q(uit)

Quits the debugger.

The next set of commands are for controlling execution:

n(ext)

Executes code until the next line of the current function is reached.

s(tep)

Executes the current line, stopping at the next executable line. If the current line is a call to another function, this "steps into" that other function, whereas next would not.

r(eturn)

Continues execution until the current function returns.

c(ontinue)

Continues execution.

run [args...]

Restarts the program. If args are provided, uses those as the value of sys.argv.

The commands you'll use to view values are:

a(rgs)

Prints the value of the current function's arguments.

p(rint) expression

Prints the value of *expression*. Usually this is not needed, since you can just type the expression to get its value; using p is necessary only when the expression is something that looks like a pdb command (including ones not shown here).

pp expression

Prints *expression* using pprint.pprint.

!statement

Executes *statement*. Usually the exclamation point is not necessary, since you can just type a statement to execute it; using! is necessary only when the beginning of the statement looks like a pdb command.

The expressions evaluated by p, pp, and typing an expression, as well as the statements executed with! or simply by typing them, are evaluated as if they had been in the code of the current stack frame. That means that arguments to the corresponding function and local names it has bound will be available. Typically a stack frame corresponds to a function definition, but comprehensions and exception handlers also create stack frames that will be visible in the debugger. You can change the current focus, or context, of the stack by moving up or down within it, one frame at a time, using these commands:

d(own)

Moves down to the next more recent frame

u(p)

Moves up to the next older frame

Moving around in the stack does not change its *contents*—that is, the state of the program is not changed; what it changes is the *context* in which debugging commands are executed and expressions evaluated. In particular, moving up to an older frame doesn't remove the more recent ones from the stack. Moving around the stack lets you investigate the values names have in different contexts without disturbing the state of the computation.

Another key feature of debuggers is the ability to set breakpoints. A breakpoint can be set at a particular line of code or in a specific function. When execution reaches that point in the code, the pdb command loop is entered. This is something you should eventually explore. More about this, and the rest of the pdb features not covered here, can be found in the pdb library module documentation (http://docs.python.org/py3k/ *library/pdb.html* or the equivalent on your computer).



The pdb loop operates in a different namespace from the top-level Python loop. Some names that are defined at the top level are not defined inside pdb, and names assigned inside pdb are assigned in the namespace of its current stack frame, not the top level, so the assignment will not survive exiting pdb.

The IDLE debugger

IDLE has its own debugger. You open it with the Debug command that replaces the Run Module command when the Python Shell is the current window. When you do so, you'll see a [DEBUG ON] message in the shell window. Naturally, you'll see a [DEBUG OFF] message when you select the Debug menu command again.



You can still use pdb in IDLE, either by calling pdb.set trace from your code or by calling pdb.pm in the interpreter immediately after a traceback.

The IDLE debugger works on similar principles to pdb, but it has a simple graphical user interface. There is a Stack Viewer window and a Debug Control window. It's difficult to describe exactly how best to use these, but experimentation will be rewarded. Even if you don't know quite what to do to take advantage of the contents of these windows, open them and get used to seeing what's in them. That will help you use them later, when you need them.

Tips, Traps, and Tracebacks

Tips

- Before writing complicated functions to do things with files and their contents, investigate whether command-line facilities are already available to do what you want. If they are, you can just use subprocess.call to invoke them.
- It is extremely important that you understand where Python looks for modules you want to import (including your own). Keep these points in mind:
 - —Import statements look for modules in each directory in the list sys.path. The system installation gives sys.path an initial value. Packages you obtain from outside the Python installation may also add to that path as part of their installation, as may your own programs.§
 - —When you open IDLE by double-clicking a file, the file's directory is added to the front of sys.path.
 - —Different IDEs on different platforms have different ideas about the "default directory" for creating, opening, and saving files. Pay attention and learn what yours is doing so that you don't save files in strange places—into the Python installation directories, in particular.
- If you get an error that a file your program is trying to open can't be found, first check that you have turned off the system feature that hides some file extensions, as described on page XXX. The next thing to do is use os.getcwd() to confirm that Python's current directory is what you think it is. Finally, use os.listdir('.') to see what files Python thinks are in the current directory. The Python installation includes extensive, well-written, helpfully organized, and nicely formatted documentation:
 - —You can find the documentation online at http://docs.python.org/py3k.
 - —You can open the documentation from IDLE by selecting the Python Docs command from the Help menu.
 - —On Windows, the documentation is installed in the Python installation directory as a .chm file in the doc subdirectory.
 - —On OS X, the documentation is a set of HTML files deep inside /Library/Frameworks/Python.framework; after opening it from IDLE, you can bookmark the URL in your browser.
 - —Bookmark the link to the Python library documentation in your browser. You will use that starting point far more than any other. If you can find the

[§] To modify sys.path, you must of course first import sys; even though its contents are built into Python, until you import it the names it assigns and defines are not in the namespace of the interpreter or a program you are executing from the command line. The same goes for other frequently used system modules, such as os.

- documentation directory on your computer, link to the library/index.html file it contains, so you can work offline. If you can't find it, you can link to its online location at http://docs.python.org/py3k/library/index.html.
- —Documentation can be downloaded in HTML, PDF, or text format from *http*: //docs.python.org/py3k/download.html. After downloading, you can install the files anywhere you'd like.
- —At the top- and bottom-right corners of every page of the HTML documentation are links to a module index and to a comprehensive index. The pages those link to are invaluable both for finding what you are looking for at the moment and noticing new things to explore.
- —If you are fond of browser windows with multiple tabs, it's a good idea to open tabs to the following pages in the documentation directory—or the online equivalent—and then save the set as a bookmark folder that you can open all at once (note that the first two files are in the same directory as the *library* directory, not underneath it):

modindex.html

The global module index

genindex.html

The general index

library/index.html

Despite the file's name, a table of contents for the entire library documentation

library/functions.html

Documentation of all built-in functions

library/stdtypes.html

Documentation of all built-in types, including their operations and methods

Traps

- It's easy to forget the exact name of the os.getcwd function, because it's not like its command-line equivalent, pwd.
- If you get an ImportError in the interpreter, make sure the current directory is what you think it is. Use os.getcwd() to check and os.chdir(path) to change it if necessary.
- Most of the time, the function you want will be in os.path, not os. If you get a 'module' object has no attribute error, don't assume you used the wrong function name—check that you did write os.path, not just os.

Tracebacks

Some of the errors you might encounter when using the utility modules described in this chapter include:

OSError

The functions in os raise OSError when given arguments of the wrong type, paths that aren't valid (e.g., a file that doesn't exist), or paths to files or directories that your program doesn't have the privileges necessary to read.

smtplib.SMTPConnectError

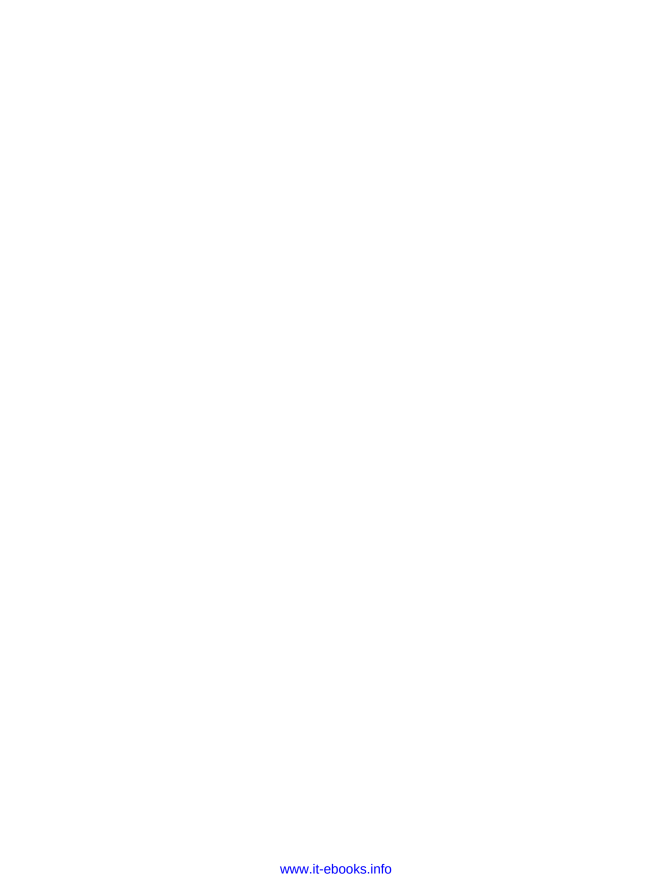
An attempt to connect to the SMTP server failed.

smtplib.SMTPAuthenticationError

Authentication with the password and username provided failed.

socket.error

A low-level communications failure occurred.



Pattern Matching

This chapter treats an enormously important topic and the Python module that supports it. We'll use its features frequently in examples later in the book. As you gain familiarity with them, you'll be using them increasingly more often. Some patience is required, though: the topic is quite technical, and you can't absorb it all at once. You'll need to come back to this periodically and explore it further. Be ambitious in experimenting with it. This is the only chapter in the book that's like this.

The mysterious topic is string pattern matching using *regular expressions*. The term "pattern matching" should be enough of a hint for you to realize how important this topic is for working with bioinformatics data. It's also important for processing web pages and for many other kinds of work a bioinformatics programmer needs to do. Among other things, restriction enzyme binding sites can be expressed as regular expressions. A carefully constructed regular expression can take the place of many lines of string manipulations, loops, and iterations. Once you've learned to use regular expressions, you'll have a very powerful tool in your hand.



If you have never encountered regular expressions before—or even if you've used only their most basic features—you should be aware that the topic is rather large, and learning it is not easy. More than most aspects of programming, learning to use regular expressions requires experimentation. Start by using the basics, and whenever you can't do what you want with what you already know, go back to this chapter (or some other documentation) and learn a little more.

Despite how much this chapter gives you to absorb, there are many advanced aspects of regular expressions that aren't covered here at all. One could write an entire book about regular expressions, and in fact several people have.* Regular expression facilities are found in the libraries of many languages, as well as in editors, IDEs, and other tools.

^{*} See, in particular, Jeffrey Friedl's *Mastering Regular Expressions*, Third Edition, and Jan Goyvaerts and Steven Levithan's *Regular Expressions Cookbook*, both published by O'Reilly.

Unfortunately, the details of regular expressions differ somewhat among different programming languages and editors, and not all resources cover Python's version exactly.



Working out the details of a regular expression you need for a particular task can be frustrating. You should design and test regular expressions outside of Python before putting them into a program. Some Python IDEs incorporate a regular expression editor and tester, but IDLE does not. An excellent interactive web page for testing regular expressions can be found at http://re-try.appspot.com (with a hyphen between "re" and "try"); you should use it often.

Regular expressions are strings used as patterns to perform matching and searching operations on other strings, which we'll call the "targets." The result of a successful operation is a *match object*, which has quite a few fields and methods providing details about the match, such as the position within the target where the pattern was found. If the match or search fails, the result is None, rather than a match object.



Python makes a distinction between matching and searching, which other languages do not. Matching looks only at the start of the target string, whereas searching looks for the pattern anywhere in the target. In discussing regular expressions it is very difficult to limit the use of the term "match" to Python's matching operations, and we will use it for both cases.

Fundamental Syntax

Within a regular expression string, certain characters have special significance. The most frequently used are similar to the wildcard characters of command-line file utilities such as 1s or dir. The wildcard characters don't have quite the same meaning in regular expressions as they do on the command line, but the idea is similar.

Backslashes have special significance within regular expressions. Backslashes also have special significance within Python strings in general. Unfortunately, what a backslash means to Python isn't always the same as what it means in a regular expression, and to include a backslash inside a regular Python string requires doubling it as \\ (just as strings can include quotes by preceding them with backslashes).

To help reduce confusion and simplify the use of backslashes in regular expression strings, Python provides a raw string syntax: preceding a string with the character r or R tells Python to ignore the usual special interpretation of backslashes within the string. Whatever backslashes appear in a raw string are just backslashes. Always use raw strings for regular expressions. Here is a demonstration of the difference:

```
>>> '\n'
'\n'
>>> len('\n')
```

```
>>> r'\n'
'\\n'
                                 # a slash followed by a newline character
>>> len(r'\n')
```

Fixed-Length Matching

Except for characters that have special interpretations, each character in a regular expression matches only that character in the target. We'll start by looking at how restriction enzyme recognition sites can be represented as simple regular expressions.

Literal matches

The recognition site of the restriction enzyme EcoRI is GAATTC. The regular expression 'GAATTC' will match exactly those characters. Since 'GAATTC' has no special regular expression characters in it, searching for it isn't really any different from using target.find(pattern). Table 7-1 gives a few examples.

Table 7-1. Literal regular expression matches

Pattern	Target	Match position	Search position
GAATTC	GAATTC	0	0
	TTGAATTC	None	2
	AATGTGAATTCT	None	5

Many restriction enzymes recognize sites with more flexibility than a literal match. For instance, EcoRII recognizes either CCAGG or CCTGG. We don't really need regular expressions to look for any of several substrings: we could easily write functions for matching or searching that do this, as in Example 7-1.

Example 7-1. Functions for multiple match and search

```
def multi match(target, patterns):
    """Return True if target begins with any of the strings in patterns"""
    for pattern in patterns:
        if target.startswith(pattern):
             return True
def multi search(target, patterns):
    """Return the first position in target where any of the strings in patterns is found"""
    return min([target .find(pattern)
                 for pattern in patterns
                 if target.find(pattern) >= 0])
```

Then we could write this:

```
multi match(sequence, ('CCAGG', 'CCTGG'))
```

to see if sequence begins with either of EcoRII's recognition sites, and this:

```
multi search(sequence, ('CCAGG', 'CCTGG'))
```

to find the first position matched by either site.

The enzyme DsaI recognizes any of CCGCGG, CCGTGG, CCACGG, or CCATGG. To use these functions for this enzyme's recognition sites we would have to provide a list of four patterns. SecI recognizes any site of the form CCNNGG; that means we'd have to include 16 sequences in the list of patterns for those sites. This would get out of hand pretty quickly. For instance, CjuI recognizes CA, followed by C or T, followed by any five bases, followed by a G or an A, followed by TG. This is $2^44^44^44^4 = 4,096$ possibilities!

Character sets

The simplest tool provided by regular expressions to manage this "combinatorial explosion" of possibilities is the ability to specify a *character set* rather than a single character. A character set consists of one or more characters enclosed in square brackets. Each character set in the pattern corresponds to exactly one character in the target: if none of the characters in the set matches that character in the target, the pattern is not a match. A regular expression for DsaI's site would be:

```
'CC[GA][TC]GG'
```

SecI's sites can be represented as:

```
'CC[TCAG][TCAG]GG'
```

And CjuI's can be represented as:

```
'CA[CT][TCAG][TCAG][TCAG][TCAG][GA]GG'
```

Character sets can contain character ranges in addition to individual characters. A range consists of two characters separated by a minus sign (-). This is equivalent to including the first character, the last character, and every character in between in the square brackets. For example, '[A-Za-z]' would match an underscore or any uppercase or lowercase ASCII letter.

Characters that have special meanings in other regular expression contexts do not have special meanings within square brackets. The only character with a special meaning inside square brackets is a ^, and then only if it is the first character after the left (opening) bracket. In that case it negates the character set, meaning the set matches any character *except* the ones specified inside the square brackets.

If you want to include a right bracket in a character set, you have to make it the first character; otherwise, it will be interpreted as the end of the character set. Similarly, if you want to include a minus sign in the character set you have to make it the first character, or it will be interpreted as showing a range.

You can also preface a ^,], or - with a backslash. While backslashes are used for a lot of things in regular expressions, they are always used with at least one other character, and technically are not one of the characters with "special meanings." To include a backslash in a regular expression—in a character set or by itself—you must use two

backslashes. The usual backslashed characters, such as \t for tab, can also be included in character sets.

Within a regular expression, a period matches any character. In effect, it represents a universal character set. Table 7-2 gives some examples.

Table 7-2. Character sets in regular expressions

Pattern	Matches
[ACTG]	One DNA base character
[A-Za-z_]	One underscore or letter
[^0-9]	Any character except a digit
[-+/*^]	Any of $+$, $-$, $/$, $*$, $^$; $^$ does not negate the others because it is not the first character in the set
[0-9\t]	A tab or a digit
•	Any character

Character classes

For convenience, the regular expression syntax offers some notation for *character* classes. These are similar to character sets in that they represent any of a certain group of characters. In fact, a character class can even be included in a character set, the way a range can. Character classes are simply convenient abbreviations for commonly used character sets and ranges that might be included as part of character sets. Table 7-3 lists Python's regular expression character classes.

Table 7-3. Character classes in regular expressions

Character	Matches
\d	Any digit
\ D	Any nondigit
\s	Any whitespace character
\ S	Any nonwhitespace character
\w	Any character considered part of a word
\W	Any character not considered part of a word

Boundaries

We've already seen in previous chapters that Python makes it very easy to treat a text file as a collection of lines. Several of the chapters that follow this one include examples of regular expressions used in processing plain text. These will include sequence data, protein data, web pages, and text in structured formats.

There is regular expression notation to indicate the beginning or end of a target, the beginning or end of a line within a target, and word boundaries. These boundary

indicators are a bit unusual in that they don't actually match any characters: they just indicate positions within the target. They're listed in Table 7-4.

Table 7-4. Boundaries in regular expressions

Character	Matches
۸	The start of a line or the beginning of the pattern
\$	The end of a line or the end of the pattern
\ A	The start of the pattern only
\Z	The end of the pattern only
\ b	The boundary between a word and nonword character or vice versa
\B	Anywhere except the boundary between a word and nonword character or vice versa

Table 7-5 gives a few examples of how boundary indicators are used in regular expressions.

Table 7-5. Line beginnings and endings in regular expressions

Pattern	Matches
^CG	A target line or string starting with CG
TATATA\$	A target line or string ending with TATATA

Variable-Length Matching

The real power of regular expressions comes from the ability to specify that certain parts of a pattern can be repeated to match the target.

Repetition

Repetition is specified using the notation shown in Table 7-6. Each of these repetition notations refers to the regular expression that directly precedes it. A single letter by itself is a regular expression, as are a character set and a period.

Table 7-6. Repetition characters in regular expressions

Character	Matches
?	Zero or one repetitions of the preceding regular expression
*	Zero or more repetitions of the preceding regular expression
+	One or more repetitions of the preceding regular expression
{n}	Exactly n repetitions of the preceding regular expression
$\{m,n\}$	Between $\it m$ and $\it n$ (inclusive) repetitions of the preceding regular expression

Using the brace notation, we can collapse the repeated elements of recognition sites into very brief expressions. SecI's sites can be represented as 'CC[TCAG]{2}GG', CjuI's 4,096 recognition sequences can be represented as 'CA[CT][TCAG]{5}[GA]GG', and the sequences XcmI recognizes can be expressed as 'CCA[TCAG]{9}TGG'—that's 262,144 possibilities represented as a 15-character regular expression!

Restriction enzyme recognition sites are fixed-length sequences. Variation in what sequences a particular enzyme recognizes arises from positions that can match two, three, or four bases instead of just one. While the number of possible combinations within a fixed-length sequence might be enormous, the number of bases to be matched is fixed for any restriction enzyme.

However, regular expressions are far more general than that. Except for $\{m\}$, the repetition characters produce regular expressions that can match targets of different lengths. We'll need more general kinds of targets to match against, so for now we'll use arbitrary base and amino acid sequences. Table 7-7 gives some examples.

Table 7-7.	Repetition	characters	in regula	r expressions
	- to F = 1111 = 11			. errp . eeeree

Pattern	Matches
CC[TCAG]{2}GG	CC, followed by any two DNA bases, followed by GG
(TA){3,8}	Between three and eight repetitions of TA, inclusive
[GC]*	Zero or more Gs and Cs (in any combination)
A+	One or more As
AT?AA	AAA or ATAA only

Greedy Versus Nongreedy Matching

The repetition-matching characters listed in Table 7-7 match the maximum possible portion of the target, provided the rest of the regular expression matches the rest of the target. This is often referred to as greedy matching. But much of the time—perhaps most of the time in bioinformatics—greedy matches absorb a far larger portion of the target than was intended.

Suppose you want to search an RNA string for the first occurrence of the polyadenylation signal AAUAAA. The regular expression r'[UCAG]+AAUAAA[UCAG]+' will match from the beginning of the string to the *last* occurrence of AAUAAA. That is because the initial [UCAG]+ grabs as much of the string as it can, which, since it includes all possible bases, is nearly the entire sequence. All that remains following that part of the match is the last occurrence of AAUAAA and one or more bases after it.

Adding a question mark after one of *, +, or ? changes the behavior to minimal possible matching, or nongreedy matching. The nongreedy matching characters are listed in Table 7-8. To return to the example of trying to find the first AAUAAA in an RNA sequence, adding a question mark after each of the plus signs in the pattern will change it so that it matches the first occurrence instead of the last. The pattern would be r'[UCAG]+?AAUAAA[UCAG]+?'.

Table 7-8. Nongreedy regular expression repetition characters

Characters	Matches
*?	Zero or more repetitions of the preceding regular expression, nongreedily
+?	One or more repetitions of the preceding regular expression, nongreedily
??	Zero or one repetitions of the preceding regular expression, nongreedily (rare)

The concept of nongreedy matching is a confusing one. When a pattern that contains repetition characters is matched against a target string, there are often many different ways the pattern can match. For example, the pattern r'[TA]+[UTAG]* could match the target 'TATATATA' by matching [TA]+ against all four TAs and [UTAG]* against no characters, [TA]* against three TAs and [UTAG]* against the final TA, and so on.

The definition of greedy matching is that each variable portion of the pattern will match against as much of the target as it can, as long as the rest of the pattern can match the rest of the string. In nongreedy matching, each variable portion of the pattern will match as little of the target as possible as long as the rest of the pattern can match the rest of the string.

Grouping and Disjunction

A portion of a regular expression can be enclosed inside parentheses. (To use (or) as a literal, either precede it with a backslash or put it inside square brackets.) This has much the same effect as using parentheses in algebraic expressions. A repetition character that follows a parenthesized regular expression indicates the repetition of the expression inside the parentheses.

Parentheses do more than just group parts of a regular expression, though. The parts of the target that match groups in a regular expression are assigned numbers in sequence (starting with 1). One part of a regular expression can refer to an earlier part using the syntax $\setminus i$, where i is the group number. Suppose you wanted to match a date in the format YYYY-MM-DD, but allow any of several characters to be used in place of the hyphen as long as the same character is used in both places. Here's one way to work this out:

```
'\d\d\d\d-\d\d-\d\d'
                                               # brute-force match using hyphen
'\d{4}-\d{2}-\d{2}'
                                               # clearer
1 '\d{4}[-.,:']\d{2}[-.,:']\d{2}'
                                               # various punctuation, can be different
2 '\d{4}([-.,:'])\d{2}\1\d{2}'
                                               # punctuation must be the same
```



Remember that when including a hyphen in a character set, it must be the first character; otherwise, it is interpreted as indicating a range.

The last step in the preceding sequence enclosed the first character set in parentheses. This allowed a later "back reference" to whatever the set matched in the target, designated by $\1$.

Groups have a much more important use than back references, though: after a match has succeeded, the characters of the target that correspond to each group can be obtained from the match object returned by functions and methods that perform matches. This is a very powerful capability, which we'll explore as we go along—not just in the rest of this chapter, but as we use regular expressions in subsequent chapters as well.

The character | between two regular expressions is similar to or in Boolean expressions. It says to try a match with the first expression and then, if it fails, try the second. Any number of regular expressions can be separated by vertical bars. Each one is tried in turn until one matches or there are none left to try. If you enclose the whole sequence inside parentheses, later parts of the same regular expression and the resulting match object can tell what part of the target matched that group without even knowing which of the alternatives was matched.

The Actions of the re Module

The re module provides functions and methods that use regular expressions for a variety of actions. Matching and searching functions and methods return match objects when successful; otherwise, they return None. We'll discuss match objects a bit later. For now, we can just treat them as true values.

Functions

First, we'll look at some of the re module's functions (the meaning of the flags arguments to these functions is described in the next section):

re.match(pattern, target[, flags])

Returns a match object if the regular expression pattern matches zero or more characters starting at the beginning of the target string, under the control of the flags value

re.search(pattern, target[, flags])

Returns a match object for the first place in the *target* string that *pattern* matches

re.findall(pattern, target[, flags])

Returns a list of all nonoverlapping matches in *target* as a list of strings or, if the pattern included groups, a list of lists of strings

re.finditer(pattern, target[, flags=0])

Returns an iterator that produces a match object for each place in target that pattern matches

re.split(pattern, target[, maxsplit=0[, flags=0]])

Returns a list of strings obtained by splitting the target at each place where pattern matches, up to maxsplit splits (all if o or omitted)

re.sub(pattern, replacement, target[, count=0[, flags=0]])

Returns the string obtained by replacing each match of pattern to target with replacement, up to count matches (all if o or omitted)

re.subn(pattern, replacement, target[, count=0[, flags=0]])

Performs the same actions as re.sub, but returns a tuple (newstring, number), where newstring is the string with the replacements made and number is the number of substitutions made

The module also provides two important functions that don't do matches or searches:

re.escape(string)

Returns a string with all characters other than letters and digits preceded by slashes, for when you want to use a string as a literal regular expression pattern and don't want any characters taking on special meanings

re.compile(pattern[, flags])

Returns a regular expression object constructed from pattern and flags; the significance of this function will be discussed shortly

Flags

Some details of matching and searching are controlled by flags. Table 7-9 describes the re module's flag values. You can refer to a flag by either its short or its long form.

Table 7-9. Regular expression match and search flags

Long form	Short form	Effect
re.ASCII	re.A	Restrict \w , \W , \b , \B , \S , and \S to matching based on ASCII characters only, not the full Unicode character set
re.IGNORECASE	re.I	Ignore case
re.LOCALE	re.L	Make \w , \W , \b , \B , \s , and \S dependent on the local language
re.DOTALL	re.S	Allow . (dot) to match a newline
re.MULTILINE	re.M	Allow $^{\wedge}$ to match at the beginning of each line in addition to the beginning of the string and $\$$ to match at the end of each line (after the newline) in addition to the end of the string
re.VERBOSE	re.X	Permit writing of formatted regular expressions that can even include comments

Functions in the module that take flag arguments expect a single value, not an arbitrary number of flags or a sequence of flags. When you need to specify multiple flags, you must use the | operator to combine them. The | operator—in an algebraic context, not within a regular expression—performs a bitwise logical or. Instead of treating the operands as two values that are each either true or false, bitwise logical operations treat each bit of each operand as a separate true or false value. For example, the bitwise or of 12 and 6 is 14:

```
1100
0110
```

It is a common computing technique to define a set of flags that are each integers with just a single bit. That allows the bitwise or operator to act as a set union operator for the flags, so that any combination of flags can be passed as a single value. For example, re. I is 2 (binary 00010), re. M is 8 (binary 01000), and RE. S is 16 (binary 10000), so if you wanted to pass the combination of the three flags as an argument you would pass the expression re.I | re.M | re.S. (The order doesn't matter.) The binary result of that expression is 11010, but using the names for the flags and the | operator allows you to ignore all that.

You probably won't use the LOCALE flag much. The ASCII flag introduces a limitation that is usually not necessary, except perhaps to avoid confusion in your mind about which Unicode characters are "space" or "word" characters. (This is an intricate topic that you'll probably want to stay away from if you can.) The IGNORECASE flag, as it suggests, ignores the case of letters in the target when performing the match.

You will use DOTALL and MULTILINE quite frequently, especially when using regular expressions to process sequence data. Without DOTALL, periods won't match newlines. Without MULTILINE, ^ and \$ will match only at the beginning and end of the target, respectively. When matching against a target string with internal newline characters, such as a sequence read directly from a FASTA file, you need these flags (specified as re.DOTALL | re.MULTILINE or re.S | re.M). Matching text containing URLs raises the same problem—you can write a regular expression to match hyperlinks (and we will), but there's no requirement in HTML that they appear entirely on one line.

The VERBOSE flag is quite different from the others. It doesn't control specific details of the match, but rather lets you write regular expressions in a more readable format, much like regular code. Unless preceded by a backslash, whitespace and comment characters are ignored.

Example 7-2 shows a regular expression that could be used to match decimal numbers. Note that the space after the d on the first line is ignored: all whitespace is ignored unless preceded by a backslash. Without the VERBOSE flag set, and assuming therefore that the comments and newlines are removed, the beginning of the pattern would read "one digit followed by one or more spaces" instead of "one or more digits."

```
r"""\d + # match one or more digits (remember to use raw strings!)
    \. # match a decimal point (not any character, because backslashed)
   \d * # match 0 or more digits"""
```

Methods

The re.compile function returns a regular expression object that encapsulates a data structure built from the pattern and controlled by the flags provided as arguments. Regular expression objects have methods corresponding to the module's match and search functions. Some of them have startpos and endpos arguments that provide more control over matches and searches than the corresponding module functions. Using the compiled version is more efficient when the regular expression is used more than once, since otherwise the regular expression will have to be compiled each time it is used.

The parameters of match object methods are different from those of the corresponding module functions. Since the pattern and flags are incorporated into the regular expression object, they aren't included in the parameters. Compiled regular expression objects have the following methods:

```
match(target[, startpos[, endpos]])
```

Returns a match object if the regular expression matches zero or more characters starting at the beginning of target

```
search(target[, startpos[, endpos]])
```

Returns a match object for the first match in the *target* string

```
findall(target[, startpos[, endpos]])
```

Returns a list of all nonoverlapping matches in target as a list of strings or, if the pattern included groups, a list of lists of strings

```
finditer(target[, startpos[, endpos]])
```

Returns an iterator that produces a match object for each match

```
split(target[, maxsplit=0])
```

Returns a list of strings obtained by splitting the target at each match, up to maxsplit splits (all if o or omitted)

```
sub(replacement, target[, count=0])
```

Returns the string obtained by replacing each match with replacement, up to count matches (all if 0 or omitted); replacement can be a function, in which case it is called with the result of each match and its return value is used to replace the corresponding match in target

```
subn(replacement, target[, count=0])
```

Same as re.sub, but returns not just the string, but a tuple (*string*, *number*) where number is the number of matches; replacement can be a function, in which case it is called with the result of each match and its return value is used to replace the corresponding match in target

Results of re Functions and Methods

Many of the re module's functions and regular expression object methods return match objects. These provide a wealth of information about the successful matches, some accessible as fields and some by calling methods.

The most powerful feature of match objects is that they capture the parts of each match in a target string that corresponds to one of the pattern's parenthesized groups. As a result, regular expressions and the functions and methods that use them can give much more than a yes or no answer to matches and searches. The groups captured in the match objects are an analysis of the structure of the matched string. We'll see how this works in examples later in this chapter.

Match Object Fields

Match object fields are accessed directly using dot notation. That is, if mobj is a match object, the startpos field will be accessed as mobj. startpos. The fields include:

startpos

The value of startpos passed to the function or method that returned the match object

endpos

The value of endpos passed to the function or method that returned the match object

rρ

The regular expression passed to the function or method that returned the match object

string

The target passed to the function or method that returned the match object

The index (integer) of the last group in the match, or None if there were no groups

Match Object Methods

Match object methods include the following. In any match object method with group numbers as parameters, 0 refers to the entire match, 1 to the first parenthesized group in the match, 2 to the second, and so on:

group([groupnumber1 [, groupnumber2 [,]]])

With no arguments, returns the entire match; with one argument returns the string that matched the corresponding group number, and with multiple arguments returns a tuple with the strings corresponding to the specified groups

groups([default])

Returns a tuple containing all the groups of the match; for any group in the pattern that was not used in matching the target, the value in the tuple will be *default* (or None if not specified)

start([groupnumber])

Returns the starting position of the target string that was matched by the group with index *groupnumber*; with no arguments, returns the whole part of the target that was matched

end([groupnumber])

Returns the ending position of the target string that was matched by the group with index *qroupnumber*; with no arguments, returns the whole part of the target that was matched

span([groupnumber])

Returns the pair (*startpos*, *endpos*) corresponding to the range of the target string that was matched by the group with index *qroupnumber*; with no arguments, returns the first and last position of the whole part of the target that was matched

expand(string)

Performs "backslash substitution" like the regular expression object sub method, but also replaces each occurrence of \1, \2, etc., in *string* with the value of the corresponding matched group

Putting It All Together: Examples

Now that we've looked at the features of the re module in some detail, it's time to put them together and see how they work in some useful examples.

Some Quick Examples

We'll start with a few simple examples to give you a more concrete idea of how to use the re module's facilities; then we'll move on to more substantial examples.

Using regular expressions to ignore case

The find and index methods of str do not provide any way to search while ignoring letter case. In some of our examples, we got around that problem by converting the target string to all uppercase or all lowercase. This is fine for simple cases, but it's awkward for situations in which you want to obtain information from the target string in its original case: you'd have to first search a single-case copy for the positions where

what you are looking for are found, then use those positions to extract the corresponding substrings from the original.

Regular expression searching provides a simple solution: just use the re.IGNORE flag. Despite their general power, there is no reason to reserve regular expressions for complicated situations. If you want to search target for string, ignoring case, this expression is all you need:

```
re.search(string, target, re.IGNORE)
```

Listing files in a directory, excluding some

Chapter 6 showed a function for listing the contents of a directory while excluding files with certain extensions. Example 7-3 shows a more flexible version of this function based on a list of file patterns instead of just file extensions. The function joins regular expressions in the list into a single pattern with the disjunction character | separating them.

Example 7-3. Directory listing filtered by regular expressions

```
def ls(path = ".", ignorepats = (r'.+\.pyc$', r'.+~$', r'^\#')):
    if ignorepats:
        pat = re.compile('|'.join(ignorepats))
        # construct an RE disjunction containing each item of
        # ignorepats to create an RE that matches any of the items
    for filnam in os.listdir(path):
        if not ignorepats or not pat.search(filnam):
            print(filnam)
```

Finding open reading frames

Regular expressions are ideal for all sorts of sequence feature detection. For example, they dramatically simplify finding open reading frames when compared to the kind of loops we saw in the examples of Chapter 4. We don't even have to worry about an extra base or two left over at the end of the sequence, the way we did in the loop-based code. Here's all we need:

```
openpat = re.compile('''
    ([TCAG]{3})*?
                                  # 0 or more codons
                                  # start codon; begin match group
     ([TCAG]{3})*?
                                  # 0 or more codons
                                  # end match group
    (TAA|TGA|TAG)
                                  # a stop codon
''', re.I | re.X )
```

The two occurrences of ([TCAG]{3})*? force the match to consider three bases at a time. The pattern [TCAG]{3} is equivalent to [TCAG][TCAG][TCAG]. The first occurrence consumes codons that precede the ATG that begins the open reading frame, three at a time. The second finds the codons within the open reading frame. The parentheses around the first one are there not because we care about the result, but simply to make the *? apply to the codon characters, since regular expressions don't allow one repetition indicator to directly follow another.

It would be nice if we could just use findall to find all the matches, but that would find the ATGs in *any* reading frame—for the same sequence, it might find one in reading frame 1 and another in reading frame 3, which makes no sense. We need to specify the reading frame we want the regular expression to use, so instead of findall we'll use match, which will force the regular expression to always start its match at the beginning of the string. We still need a loop to start each match at the end of the previous one and to collect the results, though. Example 7-4 shows a function that uses the compiled regular expression to find all the open reading frames starting in a given frame.

Example 7-4. Finding open reading frames with a regular expression

```
def open_reading_frames(seq, frame=1):
    lst = []
    matchobj = openpat.match(seq, frame-1)
    while m:
        lst.append(m.group(2))
        matchobj = openpat.match(seq, matchobj.end())
    return lst
```

An even more compact version can be written as a generator, as shown in Example 7-5.

Example 7-5. Regular-expression-based open reading frame generator

```
def open_reading_frames_generator(seq, frame=1):
    matchobj = openpat.match(seq, frame-1)
    while matchobj:
        yield matchobj.group(2)
        m = openpat.match(seq, matchobj.end())
```

The expression open_reading_frames_generator(seq, frame) will return a generator, and every time next is called on that generator the next open reading frame will be returned.

Extracting Descriptions from Sequence Files

The web page at http://rebase.neb.com/rebase/rebase.seqs.html has links for viewing and downloading DNA and protein sequences for restriction enzymes. The sequences are grouped into files according to enzyme category, but the format of the files is consistent. Assume we've downloaded one of the DNA files to the file <code>rebaseseqs.txt</code>, and we want to extract the description data for each of its sequences.

The description data in these files has a rigid format:

```
>EnzymeName RecognitionSequence SequenceLength Type
```

The number of spaces separating the fields is fixed: 3, 2, and 1. Not all sequences in these files designate a RecognitionSequence field. Examples in Chapters 3 and 4 showed a variety of ways of approaching a problem like this using comprehensions, loops, and

iterations. We're going to see now how much simpler a solution using regular expressions can be. If we can define an appropriate regular expression, we can use re.findall to find all the descriptions, doing the repetitions for us that we would otherwise have to program with a comprehension, loop, or iteration. Not only that, but a regular expression with grouping will even parse the results into their fields.

The first step is to create a regular expression that matches the whole description line. To simplify things, we'll assume that the contents of the datafile have been read and named data:

```
re.findall(r'^>.*$', data, re.MULTILINE)
```

This regular expression matches target substrings characterized by:

- > at the beginning of a line
- Followed by any characters
- Until the end of the line

We use re.MULTILINE so ^ and \$ can match lines within data; otherwise, they would match only at the beginning and end of the whole string. Using this pattern with re.search returns a list of strings such as:

```
'>M.AacDam GATC 855 nt'
```

The next step is to match the fields of the data individually.

```
re.findall(r'^>[^]+ [\w]^* \d+ .+$', data, re.M)
```

When you are first learning about them, even a regular expression as basic as this one looks quite cryptic. When you encounter one, try writing down what you think it means. Just go from left to right, as you would with an algebraic expression. This one is read as:

- A > at the beginning of a line
- One or more nonspace characters, captured as a group
- Exactly three spaces
- Zero or more word characters, captured as a group
- Exactly two spaces
- One or more digits, captured as a group
- Exactly one space
- One or more characters at the end of the line, in a group
- The end of the line

You can also use the re. VERBOSE (re.X) flag and comment the expression itself. Just be sure to put backslashes in front of any spaces inside the regular expression, because with that flag all whitespace—space, tabs, newlines—is ignored, apart from whitespace indicated by escaped characters (\ \n\t). Here's how we could combine our regular expression with the explanation that follows it:

```
^>
              \# a > at the beginning of the line
([^\ ]+)
              # one or more nonspace characters, captured as a group
| | |
              # exactly three spaces
([\w]^*)
             # zero or more word characters, captured as a group
             # exactly two spaces
\ \
(\d+)
             # one or more digits, captured as a group
              # exactly one space
(.+?)$
              # one or more characters at the end of the line, in a group
```

The result returned by findall is the same as for the earlier regular expression. We take this step in developing the regular expression to make sure that we have the pieces described accurately. For instance, if there were only two spaces after the first +, there wouldn't be any matches. In the next step we simply put parentheses around the parts we are interested in so that we can extract them from the match objects as groups:

```
re.findall(r'^{(^)}([^]+) ([\w]*) (\d+) (.+$)', data, re.M)
```

The result we get now is different, and more useful: instead of a list of strings, we get back a list of tuples of strings. Each tuple contains the portion of the matched string corresponding to one of the parenthesized portions of the regular expression. For the line:

```
'>M.AacDam GATC 855 nt'
the tuple in the list of results would be:
    ('M.AacDam', 'GATC', '855', 'nt')
```

Earlier in the chapter, it was observed that regular expressions used with bioinformatics data usually contain nongreedy repetition characters. The reason we didn't need them here is that the pattern *anchored* the match to the beginning and end of a single line. Since the goal was to extract single lines and the information they contained, using greedy repetition characters did not create any problems in this case.

Extracting Entries From Sequence Files

Also in Chapters 3 and 4 were a number of examples that involved reading descriptions and/or sequences from FASTA files. They required function definitions to use string methods to search and split either individual lines or the entire file contents. As a result they were somewhat complex.

Using regular expressions to extract entries

Regular expressions make getting the next sequence from the file a much simpler task. To extract a description and all the sequence characters, the following suffices:

```
re.search(r'^>[^>]*', src).group()
```

This reads as "match the next > that starts a line through all characters that aren't a >." This takes care of the problem of needing to read the description line to find the end of the previous sequence. We'd like to do better than that, though. First we'll separate the description from the sequence, capturing two groups:

```
re.search(r'^>(.*)$([^>]+)', src, re.M).groups()
```

The plus at the end prevents the pattern from matching a (possibly partial) description line not followed by any sequence characters, which could happen at the end of the src string. The \$ serves to split the match into everything on the description line and everything else.

Then, all we have to do is use ordinary string functions to clean up each of the two groups returned. Example 7-6 shows a complete definition of next item that doesn't use subfunctions to read the fields from the file. (The cleanup code, however, is in a separate function, so those details are out of the way of the basic matching loop.) In this example, next item searches for the next part of the text that matches the compiled regular expression, then calls extract from match to process the result. If there is no match, matchobj will be None and extract from match will return None. Otherwise, it will return a tuple of the form (description, sequence). The description, found in group 1 of the match object, is broken up into a tuple by splitting at the occurrences of | and stripping the spaces from the results. The sequence, found in group 2, has its newline characters removed.

```
pat = re.compile(r'^>(.*)$([^>]+)', re.MULTILINE)
def next item(src):
    return extract from match(pat.search(src))
def extract from match(matchobj):
    return (matchobj and
                                                     # return None when match fails
```

Example 7-6. Defining next_item for FASTA with a regular expression

matchobj.group(2).replace('\n', '')),

A really powerful feature of this solution is that we can accommodate variations in the file format with only slight changes to either the pattern or the cleanup in match item.

(([field.strip() for field in matchobj.group(1).split('|')],

Keeping track of the position between calls

matchobi.end()))

Unfortunately, this definition of next item has a major problem. Regular expression operations expect their arguments to be strings (or things like strings, such as bytes and bytearrays); they do not work with open files. The src parameter of next item must therefore be a string representing the entire file contents. However, the preceding definition will return the same item each time it is called.

We can fix this problem by keeping track of where the last match ended and starting from there the next time next item is called. This is called *state maintenance*. One way to keep track of a value in between function calls is:

- 1. Add a parameter for the previous value to the function.
- 2. Return the value along with the other value(s) the function returns.
- 3. Have each function supply the previous value as one of its arguments.
- 4. Assign a name to the value part of the result of the function.

This is probably easier to illustrate with an example than with a written description. Let's consider how we could implement a findall function for plain strings using str.find. Example 7-7 shows a straightforward iterative definition like many we've already seen. Each time the target substring is found, the function adds its position to a list it is accumulating and then searches again, starting at the next character after the substring.

Example 7-7. A findall function for ordinary strings

```
def findall(string, substring):
    """Return a list of all nonoverlapping positions in string where substring appears"""
    positions = []
    pos = string.find(substring)
    while pos >= 0:
        positions.append(pos)
        pos = string.find(substring, pos + len(substring))
    return positions
```

Example 7-8 shows an alternative approach with two functions: one just finds the next occurrence of the target, and the other calls it and collects the results. There's no particular advantage to doing things this way for this simple example; it simply illustrates the technique so we can use it for something more substantial next.

Example 7-8. Keeping track of a value between calls

```
def findnext(string, substring, startpos=0):
    """Return the position of the next nonoverlapping occurrence of
    substring in string, beginning at startpos where substring appears"""
    pos = string.find(substring, startpos)
    if pos < 0:
        return -1, -1
    else:
        return pos, pos + len(substring)
def findeach(string, substring):
    positions = []
    pos, startpos = findnext(string, substring)
    while pos >= 0:
        positions.append(pos)
        pos, startpos = findnext(string, substring, startpos)
    return positions
```

The important point to note about these two functions is that findnext needs to know where to start looking for the substring each time it is called. It's up to findeach to keep track of that next starting position between calls. We've defined findnext so that it returns two positions: the position where the substring starts and the position where search should recommence. In findeach, the two values returned from findnext are assigned, and the second is fed back into findnext when it is called. This example is so simple that findeach could compute where to start the next search as easily as findnext could, but if findnext does anything more interesting that might well not be the case.

Example 7-9 shows a definition of next item that applies this technique to keep track of the search position between calls. It also shows an example of the function's use.

Example 7-9. Defining next_item to return and receive a position

```
def next item(src, pos):
    return extract from match(pat.search(src, pos))
def extract from match(matchobj):
    return ((None, -1)
                                                         # pos doesn't matter here
            if not matchobj
            (([field.strip() for field in matchobj.group(1).split('|')],
              matchobj.group(2).replace('\n', '')),
             matchobj.end()))
                                                         # new value of pos to be remembered
def find item(src, testfn):
    item, pos = next item(src, 0)
    while (item):
        if testfn(item):
            return item
        item, pos = next item(src, pos)
```

Another thing we can do is wrap next item in a generator. Keeping track of the values of parameters and other names assigned in the function is one of the primary responsibilities of the generator mechanism. Another is keeping track of where the yield occurred in the definition (there may be several) so that execution can resume right after that statement when next is called on the generator object. Example 7-10 shows what that would look like

Example 7-10. Defining next_item as a generator

```
def item generator(src):
    pos = 0
    item, pos = next item(src, pos)
    while item:
        vield item
        item, pos = next item(src, pos)
def find item(src, testfn):
    itemgen = item generator(src)
    item = next(itemgen)
```

```
while (item):
    if testfn(item):
        return item
    item = next(itemgen)
```

We've chosen to leave the definition of next item unchanged from its definition in Example 7-9. Instead of adding complexity to next item and extract from match, we've defined a second function to produce the generator object that keeps track of the position. Each time next is called on the generator object returned by item generator, execution resumes in item generator and continues until the yield. All this generator is doing is holding onto the value of pos between calls to next item.

Notice also the changes in find item. It no longer does anything to keep track of the position and provide it as an argument to next item. In fact, it no longer calls next item at all. Instead, it calls item generator once at the beginning of the function, then uses next(itemgen) each time it wants another item.



Example 5-4 showed a class whose methods share state using instance fields. The problem here is not quite the same as the one solved by the class example. There, the emphasis was on avoiding a lot of useless argument-passing for values that were used unchanged by many methods. Here, the challenge is to maintain changing state between one function call and the next, not just share it. In the technique demonstrated in the preceding example, one function returns values to its caller that it will need again later, and the caller passes them back as arguments the next time it calls the function. State maintenance is even more of a reason to use a class than state sharing, and usually you would use a class rather than the tricks of the previous example. We'll see examples of a two such classes in Chapter 8 (in Examples 8-17 and 8-18).

Buffering input

Another problem with these definitions is that code using them must read the entire contents of a file to pass find item. If the file is extremely large this could take an inconvenient amount of time and space, especially if the sequence is located early in the data. And even if the intent is to do something with every sequence in a file, you still may not want to read the entire contents all at once.

The solution to this sort of problem is to interpose a buffer between the file and next item. A buffer is a standard programming mechanism that turns processing a source or destination—reading from a file, writing to a file, searching a string, etc. into a two-step process. A file buffer contains a moderate portion of the file's bytes or characters. Reading is done from the buffer, rather than from the file itself. When the end of the buffer is reached, its contents (or most of them, anyway) are discarded and another piece of the file is read into the buffer. For writing, the process is reversed: writing is directed to the buffer, and when it's full its contents are written out to the file and the buffer is emptied. (This is called *flushing* the buffer.)

The main reason for buffering is that input and output operations that involve hardware storage devices are many orders of magnitude slower than reading from and writing into the computer's memory. Buffering replaces many small external interactions with many small internal interactions and only the occasional bulk transfer from or to external storage. As noted in Chapter 6 (in "The Python runtime environment: sys" on page 213), sys.stdout is a buffered file stream for this reason.

Python's implementation of input and output streams includes buffering capabilities, but programs can also implement their own, higher-level buffering mechanisms. We can add a buffer to Example 7-10's definition of item gen, enabling it to pass strings to re functions and methods without first having to read the entire contents of a huge file into memory. This will improve efficiency in the event that a call to find item locates a targeted item after examining only a small portion of the file.

The strategy will be to read a number of characters from the file and read sequences from the string returned; then, when all the sequences of the string have been read, another chunk of characters will be read. What's tricky about this is that the string beginning with the last > is almost definitely incomplete; the rest of the sequence will be in the next chunk.

The process is illustrated in Figures 7-1, 7-2, and 7-3. It's not easy to develop code like this, but it shouldn't be too hard to understand.

Example 7-11 shows a get item function added to manage the buffering.

Example 7-11. Buffered regular expression search of a file

```
def get item(fil, buffer, pos, chunksize):
     ""Return a possibly incomplete item along with the new value
    of buffer and the end position of the successful match"""
    item, endpos = next item(buffer, pos)
                                                                 # initialize loop
    while not item:
                                                                 # look for next item
        chunk = fil.read(chunksize)
                                                                 # read next chunk
        if not chunk:
             return None
                                                                 # end of file
                                                                 # add chunk to buffer
        buffer += chunk
        item, endpos = next item(buffer, pos)
                                                                # try again
    return item, buffer, endpos
                                                                # beginning of an item
```

The purpose of get item is to call next item until it returns an item. Each time next item fails to find an item, get item reads chunksize characters from the file and appends them to the buffer. When enough of the file has been read into the buffer for next item to find an item, get item returns the item, the (possibly extended) buffer, and the ending position of the match. This follows the approach discussed earlier, whereby values are maintained between calls by returning them to the caller and having the caller pass them back in. In this case, two values are tracked: the buffer itself and the position within the buffer where the search should resume.

The end of the buffer might contain a FASTA description and the beginning of its sequence, but not the entire sequence. How can we know whether get item has

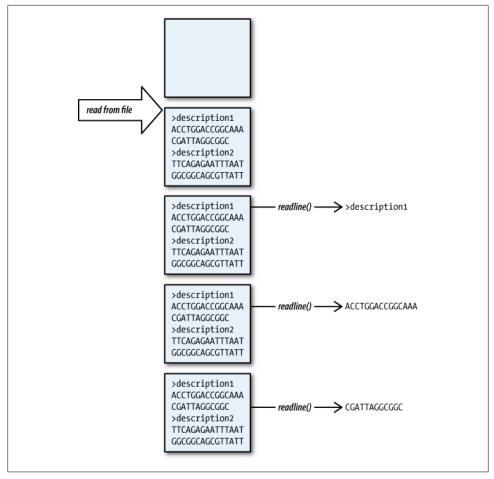


Figure 7-1. Buffering input while doing searches, part 1

returned a complete sequence? The solution is to always read a second sequence ahead of the one to be returned. As long as the beginning of another sequence is found, we know the current one is complete. When the current sequence is the last in the buffer, looking for the next one will cause the buffer to be extended until a next sequence is found. At that point, the current sequence must be reread to get its full contents.

The outline of the code is similar to a lot of loop-based functions: it reads an item, then loops until there are no more items. In this case, however, the code starts by reading two items and loops until there is no more second item. The technical term for reading one item beyond the targeted one is *lookahead*, for obvious reasons. Lookahead is required whenever some information from the next item is needed to process the current item (including, at a minimum, whether there is a next item). In the normal case, each time around the loop, the name of the first item is rebound to the second item and a new second item is read. This code also has to deal with the abnormal case when no

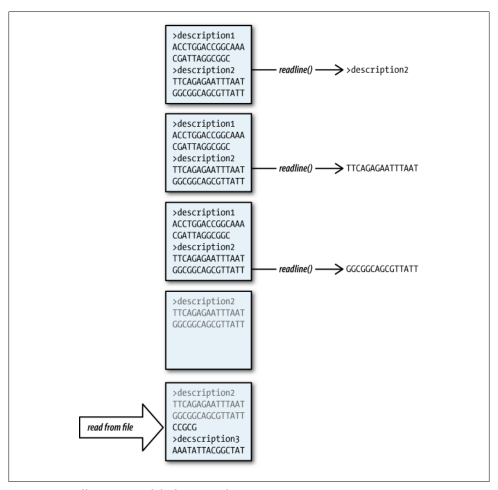


Figure 7-2. Buffering input while doing searches, part 2

second item is found, because that is when the buffer will be extended (except at the end of the file). The outline looks like this:

```
1 initialize
2 first item = read item
3 nextitem = read item
4 while nextitem:
   if finding the next item caused the buffer to be enlarged
      reread item from the buffer because it was incomplete
      shorten buffer to exclude previously read items
8 yield item
   item = nextitem
10 nextitem = read item
11 yield item
                 # at end of file
```

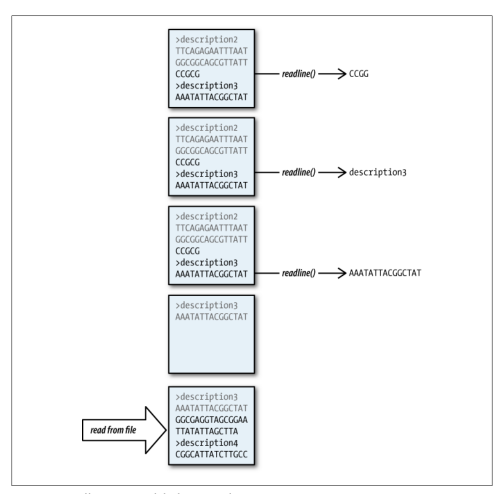


Figure 7-3. Buffering input while doing searches, part 3

Example 7-12 shows code that implements this outline. The solution is complicated by having to maintain three positions: the beginning of the current item (curpos), the beginning of the next item (nxtpos), and the end of the next item (nxtend). (At the end of the buffer, nxtend will equal the length of the buffer). A numbered line in the code, together with any following unnumbered lines, implements the correspondingly numbered part of the outline.

Example 7-12. Buffered regular expression generator for a file

```
def item_generator(file, chunksize=4000):
    curpos = 0
    itm, buffer, nxtpos = get_item(file, '', curpos, chunksize)
    nxtitm, buffer, nxtend = get_item(file, buffer, nxtpos,
```

```
4
      while nxtitm:
5
          if len(buffer) > nxtend:
6
              itm, nxtpos = next item(buffer, curpos)
              nxtend -= nxtpos
7
8
          vield itm
          itm, curpos, nxtpos = nxtitm, nxtpos, nxtend
9
           nxtitm, buffer, nxtend = get item(file, buffer, nxtpos,
10
       yield next item(buffer, pos)[0]
11
```

Tips, Traps, and Tracebacks

Tips

The importance of constructing regular expressions piece by piece cannot be emphasized enough. Make sure each pattern is working as expected before making it more complex. Does this sound familiar? It should: this book has repeatedly stressed this approach for defining functions. Unfortunately, regular expressions don't have the ability to call other regular expressions, so you can't really build them out of truly separate pieces. You can, however, still proceed in a step-by-step manner, increasing the complexity as you go.

Debugging regular expressions is far more difficult than debugging programs. You can't work your way step by step through the match the way you can with a Python debugger. There are tools for testing and debugging regular expressions, but even without one you can often accomplish a lot by simply printing the results of a findall, one to a line. Another trick that's useful when a regular expression isn't doing what you expect is to remove parts of it gradually and repeat the search until you locate the problem.

As you start building more complex regular expressions, you will experience more than a little frustration. Here are a few pointers:

- Use a regular expression testing tool such as re-try. You don't need the entire source file for testing your regular expression—just grab a section that will produce a match or two once your regular expression is working.
- Remember to use raw strings for regular expressions.
- In general, it's probably not worth trying to tune a regular expression so that a group such as FASTA sequence characters matches multiple lines but excludes the newlines. Just use replace('\n', '') on the result.
- Separating fields by a delimiter character such as a vertical bar on a FASTA description line can be done within the regular expression, but if you are having trouble accomplishing that, just have the expression match the whole line and use the following to get a list of the fields:

```
[field.strip() for field in line.split('|')]
```

• To search any of a list of patterns, search for:

```
'|'.join(list of patterns)
```

Traps

• Remember to use the | operator to combine flags; re functions that take a flag argument, including re.compile, only accept one such argument. If you forget and use commas instead you will get an error message like the following:

```
TypeError: fn() takes at most 2 positional arguments (3 given)
```

The exact numbers will depend on which function you called and how many arguments you supplied.

- Beware of vertical bars in regular expressions! They represent disjunction. If you are trying to match a vertical bar, as in FASTA headers, you must use a backslash $(\backslash |).$
- When attempting to capture a group, make sure the repetition characters are inside the group unless you really do mean to repeat the group. That is, if you want a sequence of digits, write (\d+) and not (\d)+. The latter will return the last digit only, a rather mysterious result. This is an easy mistake to make, especially if you casually put parentheses around a small part of a partially completed regular expression.
- When matching the contents between "delimiter" characters—quotes, parentheses, brackets, braces, angle brackets, etc.—a * or +, and even a nongreedy *? or +?, will often match too much of the target string. In these situations you should match a repetition of "anything but the closing delimiter." For example, to match something enclosed in quotes, use "[^"]+" instead of ".+?".
- Make sure to use the re.DOTALL (re.S) flag if a newline can occur within text you want matched by .*, .+, .*?, or .+?, and not to use it if you don't want newlines matched.
- Make sure to use re.MULTILINE (re.M) if you are using anchors—^ for the beginning of a line and \$ for the end. Otherwise, ^ will match only the beginning of the entire string, and \$ will match only the end.
- If you use re. VERBOSE (re.X), make sure to put a backslash in front of any space that you mean to be part of the pattern; otherwise, it will be ignored just like the whitespace you use for making the string readable.
- A negated character set such as [^>]* will match line feeds. This pattern:

```
r'^>[^>]*'
```

will match an entire FASTA entry—the description and all of the sequence characters, with newlines embedded, regardless of whether re.DOTALL is specified.

If re.MULTILINE is specified, a dollar sign added to the end of that expression will have no effect because the character set will still match newlines. To prevent negated character sets from matching newlines you must include the newlines explicitly in the character set, either as \n or \s (which would exclude all whitespace).

Tracebacks

• The same kinds of errors that occur when parentheses aren't balanced in code, especially in algebraic expressions, can easily happen with group parentheses in regular expressions, producing the error:

```
sre constants.error: unbalanced parenthesis
```

• The error:

```
sre constants.error: nothing to repeat
```

can be particularly irksome. If you are developing a pattern incrementally, the obvious place to look for the source of this problem is where you last added a repetition character, or perhaps a pair of parentheses.

• The error:

```
sre constants.error: multiple repeat
```

means that somewhere in the regular expression there are adjacent repetition characters. Perhaps the most likely culprit is putting the question mark before the plus sign or asterisk in a nongreedy repetition, instead of after. This problem can also arise when, while developing a regular expression, you accidentally remove the character(s) that separated two repetition characters.

• Unfortunately, the error won't point you to where in the expression the problem occurs; in fact, if you are using re.VERBOSE (re.X) it won't even show you which line to look at—the error always points to the end of the function call in which the expression appears. Note also that the line number of the offending function call appears early in the traceback, followed by many calls to compile, parse, and related subfunctions. This contrasts with most tracebacks, where it is the last few lines that show you where in your code the error occurred.



Structured Text

In Chapter 6, we took a very brief look at the csv module that is used to read and write lines of tab- or comma-separated values, with each line corresponding to one item in the file. We've also looked at a variety of ways to scan files looking for certain patterns of data, including using str methods and regular expressions. Files that are in tab- or comma-separated values format, FASTA files, GenBank files, and many other file formats encountered in bioinformatics work are called *flat files*.* What is "flat" about them is that they are just text files: the data has no explicit structure beyond agreed-on conventions regarding special characters, blank lines, whitespace, etc. They can have introductory material before the data, other material after the data, several sets of data in one file, and so on.

The opposite of "flat" in this context is *structured*. A structured text file contains *elements*, each of which can have *attributes* and/or "sub" or *child* elements. There can be different kinds of elements, and in general there are rules specifying what attributes and children each kind of element can have. The linear approaches for processing text files that we've seen so far are inadequate for structured files, essentially because the files are two-dimensional. This chapter describes some ways to process structured files.

HTML

An obvious example of a structured file format is basic HTML. (We'll ignore all the fancy stuff like JavaScript, frames, and so on.) HTML's structuring rules are a bit loose, but we can ignore those details. The top-level structure of an HTML file is:

* In computer science the term "flat file" usually has a stricter meaning, referring only to text files with one item per line, each having fields designated by separators (commas, tabs, vertical bars, spaces, etc.) or conforming to some specified number of characters. Files in formats such as FASTA and GenBank would be considered "free form," even though they have some regularity.

```
<body>
         . . . tags and text . . .
    </body>
</html>
```

Each HTML element opens with a start tag, which begins with one of the tag names defined by the HTML specifications and is enclosed in angle brackets (<>). Tags may have attributes, in the form attribute - name = attribute value. Elements may also have content—either text or other elements—that appears after the start tag. Tags with no content may be closed by a slash at the end of the start tag instead of an end tag: for instance, <hr/> to draw a horizontal line. Comments, which can span many lines and cause the text between them to be ignored by a browser displaying the HTML, begin with the characters <!-- and end with -->.

The end of a tag with content may be indicated by </tagname>, where tagname is the same as the name in the start tag, but for the most part the HTML standards don't require that. For instance, it is common practice to mark the beginning of a paragraph with a tag but not to bother with the closing , leaving the browser to figure out that the paragraph has been closed because a new one has been started (among other reasons).



People often use the term "tag" to refer to the structure element a tag specifies, in addition to using it for the tag itself. In the rest of the book we will use just the tag name when talking generically about a tag, as in the phrase "an hr tag." When we specifically mean a start or end tag, we will use angle brackets.

Some tags in a file contain others. For instance, the top-level html tag contains the body tag, which contains the rest of the tags in the file. Many of those could be tags for headers or paragraphs whose content is just text. Other kinds of tags have "child" tags as their content. Lists are an example:

```
Item 1
  Item 2
  <a href="http://www.example.org">Item 3</a>
```

This bit of HTML says to create a numbered list (o1) containing three items (each 1i).

There are other sorts of things inside HTML files, such as a few tags at the beginning that contain information about the file itself and embedded scripts that you usually won't want to consider as part of the document. There are also special notations such as named entities for punctuation, Unicode characters, and so on. † You'll see those most often where characters with meaning inside HTML itself are part of content; for

[†] You can find some good references at http://www.w3schools.com/tags/ref_entities.asp, http://www.entitycode .com/#common-content, and http://www.escapecodes.info.

example, & or & for ampersand and < or < for <. Each named entity begins with an ampersand, ends with a semicolon, and contains either letters or a pound sign followed by a base-10 number that refers to specific characters in standardized character sets.

Unicode Characters in HTML Files

HTML files often contain *actual* Unicode characters, not just named entities that *represent* them. Any attempt to read from a file that contains Unicode characters will cause an error unless you provide the optional keyword argument <code>encoding='utf-8'</code> in the call to <code>open</code>. Since the default encoding is a subset of Unicode, you can use that argument even with files that contain just ordinary ASCII characters.

If you want to use the file's actual encoding rather than guessing that 'utf-8' will work, you can find it in the file itself. Any HTML file that contains non-ASCII characters should have a tag like the following in its head section:

```
<meta http-equiv="Content-Type" content="text/html; charset=UTF-8">
```

The following code will extract the encoding from an HTML file. It opens the file in binary mode to avoid any encoding issues, reads some characters, uses a binary regular expression to look for the tag that specifies the encoding, closes the file, and (if it finds one) returns the value of the tag's encoding attribute decoded into a string. Subsequent examples that read from HTML files will use get_html_file_encoding to determine the encoding argument they should provide when opening the file for processing:

```
encoding_pat = re.compile(
   b'<meta [^>]*?content *= *[^>]*' +  # just to fit line
   b'charset *= *([a-zA-Z0-9-]+)',
   re.I | re.A)

def get_html_encoding(html):
   encoding = encoding_pat.search(html)
   return encoding and encoding.group(1).decode()

def get_html_file_encoding(filename):
   with open(filename, 'rb') as file:
        return get_html_encoding(file.read(2000))
```

Simple HTML Processing

Some simple search or replace operations can be performed on HTML files using a text editor, a web design tool, Python, or another programming language. Suppose you find a web page with many interesting hyperlinks, and you want to extract them from the rest of the page. You could just search for '<a ', then for 'href=', and so on, but there are too many possible variations to handle. You need regular expressions.

Searching HTML text

In this section we'll develop a program to extract all hyperlinks and associated text from a web page saved as an HTML file. The program will look a lot like some of the other examples we've already seen in the book. Example 8-1 shows the high-level functions that won't change as we develop the code further and the test at the end that lets us run it as a command-line program.

Example 8-1. Extracting <a> links from an HTML file, step 1

```
def print atags in files(lst):
    for filename in lst:
        print(64*'-')
                                                    # draw separator line
        print_atags in file(filename)
def print atags in file(filename):
    with open(filename, encoding=get html file encoding(filename)) as file:
        print atags(file.read())
def print atags(string):
    for atag in get all atags(string):
                                                    # defined below
        print atag(atag)
def print atag(results):
   print(textwrap.fill(results[0], 75),
         textwrap.fill(results[1], 75),
         sep='\n',
         end='\n\n'
if name == " main ":
    if len(sys.argv) < 2:</pre>
        print("Usage: show atags htmlfile ...")
    else:
        print atags in files(sys.argv[1:])
```



This time we've shown the definitions "top-down"—i.e., callers above callees. The order of the function definitions within a file doesn't matter to Python; it's up to you to decide how to organize the file for easiest understanding. Often that means putting the "high-level" functions at the top, so someone interested in using the program can see them without having to read through all the details. Then again, if you begin the program with a docstring describing the functions meant to be called from other files, there would be no need for a user to read the code to find them.

Next we'll present a series of definitions for get all tags, omitted from the preceding code. URL references span a wide range of formats. Before trying to formulate regular expressions to match them, it's well worth defining what we're trying to capture. As the program gets used it may produce some unexpected output, at which point we can choose to accommodate the new example it reveals or make a note that we are ignoring the problem.

There are a variety of start tags that can contain href attributes, such as , but we are only interested in <a> tags. If we discover that we are interested in others, we can always add them later. Even in trying to describe the URL pattern, we will follow the incremental approach exhibited in our earlier code examples. We'll start by capturing the <a> tags, their attributes, and their content and see what they look like before tackling more complex aspects of the patterns. That will actually bring us quite far toward our goal, but, as we'll see, may include a variety of extraneous information. Example 8-2 shows the initial regular expression.

Example 8-2. Extracting <a> links from an HTML file, step 2

Here are some of the results extracted from the output of the program run on the results of an advanced search for the phrase "nonsense mediated decay" at http://biology.plosjournals.org. (The actual output would be several hundred lines long.) For each link the content is printed first, followed by the attributes of the <a> tag starting on a new line. Where more than two lines are shown it is because one or both of these lines overflowed the line length in the Python window where the code was executed:

```
PLoS.org
href="http://www.plos.org" title="PLoS.org"

Site Map
href="http://journals.plos.org/plosbiology/sitemap.php"
title="PLoS Biology Site Map"

Readers' Responses
href="http://biology.plosjournals.org/perlserv/?request=read-response&
issn=1545-7885&past_days=360" title="For Readers: Readers' Responses"

A Combinatorial Code for Splicing Silencing: UAGG and GGGG Motifs
href="http://biology.plosjournals.org/perlserv/?request=get-document&
doi=10.1371/journal.pbio.0030158"

Posttranscriptional Gene Regulation by Spatial Rearrangement of the 3′
Untranslated Region
href="http://biology.plosjournals.org/perlserv/?request=get-document&
doi=10.1371/journal.pbio.0060092"
```

Now that we have the attributes and the content, we want to do some cleanup. First, we'll extend the pattern to extract the value of the href attribute. This is a bit tricky the equals sign can be surrounded by spaces and newline characters, and the value can be surrounded by a pair of single or double quotes. We'll allow for the quotes, but we won't worry about them matching, and quotes aren't allowed inside the value of the href attribute. We will also assume there is no whitespace within the value of href. The modified regular expression is:

```
r'''<a.*?href\s*=\s* # href=, optional spaces
       ['"]? # optional quotes
[(^'"\s]*) # the value: no ', ", or whitespace
['"]? # optional quotes
[^\]*? # the rest of the tag
(.*?) # capturing the entire content
 </a>
                                           # the </a> end tag
```

The results for the <a> tags shown previously are now:

```
PLoS.org
http://www.plos.org
http://journals.plos.org/plosbiology/sitemap.php
Readers' Responses
http://biology.plosjournals.org/perlserv/?request=read-response&
issn=1545-7885&past days=360
A Combinatorial Code for Splicing Silencing: UAGG and GGGG Motifs
http://biology.plosjournals.org/perlserv/?request=get-document&
doi=10.1371/journal.pbio.0030158
Posttranscriptional Gene Regulation by Spatial Rearrangement of the 3&#8242:
Untranslated Region
http://biology.plosjournals.org/perlserv/?request=get-document&
doi=10.1371/journal.pbio.0060092
```

For the next step, we'll replace some HTML entities with their ordinary equivalents. This is a perfect use for a dictionary, a small portion of which is shown along with a function that uses it and a changed version of print atag in Example 8-3. (The regular expression doesn't change.) With this code, the ′ in the last title in the previous output is replaced with a single quote.

Example 8-3. Extracting <a> links from an HTML file, step 3

```
html entities = {
# Reserved characters in HTML
       '"':'", '"':'"'
'&':"", ''':""
'&':'&', '&':'&',
'<':'<', '&lt;':'<',
'&#62;':'>', '&gt;':'>',
```

```
# ISO 8859-1 symbols 160-191, 215, 247
    ' ' : ' ', ' ' : ' ',
# Math symbols
    '–' : '-', '–' : '--',
    '′' : "'', '′' : "'',
    '″' : "'', '′' : "'',
}

def replace_substrings(string, dictionary):
    """Return a copy of string in which every key of dictionary has been replaced with the corresponding value, in arbitrary order"""
    for key, value in dictionary.items():
        string = string.replace(key, value)
    return string
```

Although they didn't happen to show up in the one web page used for the preceding examples, experimentation with other web pages revealed that sometimes the value of an href attribute begins with a pound sign (#). These are document fragment markers used at the end of a URL to specify a location within the page containing the marker. We don't need these, so we want to exclude <a> tags whose href attributes have a value beginning with a pound sign.

We could exclude these tags by adding \# to the list of forbidden characters where the value of the href attribute is captured. (The backslash is required to prevent Python from interpreting the # as the beginning of a comment.) The problem is that although this would make the value match fail, it wouldn't make the overall match fail: all we'd get is an empty href value with whatever content the fragment identifier was associated with.

Nothing says we have to do all the work with a single regular expression, though. We can use one regular expression to extract something from the results obtained using another one, or we can filter the results we get back from a call to findall. To solve this problem we'll filter the results, as shown in Example 8-4.

Example 8-4. Extract <a> links from an HTML file, step 4



The change is in the function <code>get_all_atags</code>, and it justifies having made this a separate function. When this was added during development of the earlier stages, it seemed entirely superfluous; there was no indication it would ever do anything other than a <code>findall</code>. It was placed there on principle, but now it turns out to have been a good decision. Note how this works—it's important.

The code should now give us what we want from the value of the href attribute in the <a>> tag. Some of the values will be ugly because they form HTTP requests with arguments, as indicated by the presence of a ? in the URL, and some of the request strings are quite long. For example, the URL corresponding to a link to "next" on the page with the PLoS search results looks like this:

```
?request=advanced-search&search fulltext=1&row start=11&
issn=15457885&surname type=all&surname=&fname type=all&fname=&aff type=all&
aff=&fundsrc type=all&fundsrc=&anywhere type=phrase&
anywhere=nonsense+mediated+decay&title_type=all&title=&abstract type=all&
abstract=&biblist type=all&biblist=&tblfig type=all&tblfig=&
jrn issn=15457885&subj id=all&vol pubdate start=&iss pubdate start=&
vol pubdate end=&iss pubdate end=&vol no start=&iss no start=&vol no end=&
iss no end=&limit=10&order=score&document count=12#results
```

We could filter these, or perhaps give the program's user that choice, but since so many web pages offering document retrieval do so by way of requests rather than URLs linking to documents directly, we'll leave them in. That will allow the results to be used as real URLs, perhaps in an HTML page the code writes.

There's one other problem with this code that we'll put off dealing with until the next chapter: many URLs on a page will not begin with a site address (or a protocol). These are addresses relative to either the root of the site from which the page was retrieved or the directory from which the page was retrieved (the former is the case if the partial URL begins with a slash). For now, we won't be able to extract sufficient information from these URLs for them to be used to retrieve a page or submit a request.

Next, we'll turn our attention to the content extracted. The challenge here is that the content can contain all sorts of markup, and we want the output to be plain text. This is another case where it is too difficult to insert this additional processing into the existing regular expression. Instead, we'll use another one to remove the tags from the content and add that step to get all atags. We'll also add a test in print atags to avoid printing empty content. The comprehension in get all tags would be a little messy if we didn't unpack the results of findall, so we add that in Example 8-5 too.

```
Example 8-5. Extracting <a> links from an HTML file, step 5
tagpat = re.compile(r'' < [^>]+?>'') # easy - anything within a pair of <>s
def remove tags(content):
    return tagpat.sub('', content)
def get all atags(string):
    return [(url, remove tags(content))
            for url, content in atagpat.findall(string)
            if url[0] != '#']
def print atags(string):
    for atag in get all atags(string):
        if atag[1]:
            print atag(atag)
```

More elaborate HTML pattern matching

Example 8-6 shows a simple program for printing an outline of a web page based on its heading tags (h1, h2, ..., h6). (Most modern web pages use other organizational techniques, so you might not find many—or any—heading tags on a page you download, but heading tags are still used for simple pages.) Try this function with the path to a simple web page with header tags that you've saved to the disk.

Example 8-6. Extracting an outline from an HTML file

```
pattern = r'<h(\d).*?>(.+?)</h'
def print_outline(filename):
    with open(filename) as fil:
        for level, hd in re.findall(pattern, fil.read()):
            # indent for each level and print the level in brackets
            print("{0}[{1}] {2}".format(' '*3*int(level), level, hd))</pre>
```



When running Python from the command line, you may find that attempts to print Unicode characters cause errors. If so, this is because your terminal application doesn't support Unicode. In Unix environments, if you set the environment variable LANG to something like en_Us.UTF-8 before running Python you will be able to print Unicode characters. You won't run into this problem in IDLE, though, because its shell is built on top of code that supports Unicode.

Even if you can find a page that uses header tags for its general organization, there may be other markup around the header contents that ruins the output. For instance, go to http://www.ncbi.nlm.nih.gov and save the BLAST QuickStart document bookshelf/ br.fcgi?book=comgen&part=blast to your disk. Then load the definition of print_out line and call it on the name of the file you used when you saved the document.

The output will be an outline, but it contains extraneous formatting detail—in particular, a lot of span tags. (span tags define formatting attributes for the content they surround.) Here's a sample line from that file:

```
<h4> <span class="mw-headline"> Documentation </span></h4>
```

We can filter out the span tags with a more complex regular expression, as shown in Example 8-7.

Example 8-7. Filtering span tags from HTML

```
</h
                 # header end tag (we aren't checking that this ends
                 # the one just started)
                        re.IGNORECASE | re.DOTALL | re.VERBOSE)
def print outline(filename):
    with open(filename) as fil:
         for level, pretag, optstart, posttag, optclose in \
             pattern.findall(fil.read()):
print("{0}{1}{2}".format(' '*3*(int(level) - 1), pretag, posttag))
```

With some practice, you'll be able to do this sort of thing yourself. However, unless you have a lot of web pages to process that use tags in a similar way, your attempts to cope with all the details by writing increasingly complex code and regular expressions won't be worth the effort and frustration. Instead, either use several regular expressions or a mix of regular expressions and regular programming techniques.

Turning HTML into plain text

One important use for str.replace is to delete all occurrences of a substring from a string, by providing an empty string as the replacement. That is also an important use of re.sub. As a simple (though not entirely realistic) example, let's turn an HTML file into plain text by removing all the tags. The results will be pretty much a mess, since they'll reflect the spacing and line breaks of the original HTML rather than any kind of formatted text equivalent. More useful results could be obtained if the patterns were tuned to specific kinds of HTML pages.

Before we remove the tags we should skip to the **body** tag that starts the page's content, so we don't have to deal with the complexities of what can appear before that. There is also the complication that pages can contain script code placed between a <script> start tag and a </script> end tag. Although usually these occur in the <head>, they can occur anywhere in the **body** too. There can also be comment tags, which can get tangled up in the other tags. (Sound familiar? This is another "skip introduction.") Our code will proceed in these steps:

- 1. Skip to the **<body>** tag.
- 2. Remove all <script> ... </script> sections of the body.
- 3. Remove all comments (<!-- ... -->).
- 4. Remove all other tags.

We'll use re.search with the re.IGNORECASE flag to locate the <body>, <script>, and </script> strings, as just described. Normally we would define a separate function for each of these steps, but since each one after the first would do the same thing—one call to sub—we can do something a bit more clever: we can just loop through the patterns in succession in one function. This makes it easy to extend the program. For instance, if we also want to remove the contents of forms or all lines with just spaces, we can add another pattern for that to the patterns list. Example 8-8 demonstrates.

Example 8-8. Extracting plain text from HTML



HTML pattern matching is notoriously tricky. In HTML patterns, even nongreedy matches often consume too many of the target string's characters. If you look back at the regular expression patterns in the preceding examples, you'll see that there are a lot of places where a character set is used to *exclude* one or two characters. The excluded characters appear right after the repetition characters (for example, [^>]*?>, which matches everything up to the first >). Typically it is angle brackets that govern the structure of the match, but quotes and other delimiting characters play an important role too. If you are having trouble with a pattern that matches too much, try using this technique to restrict the repetitions it contains.

Structured HTML Processing

The techniques and examples we've seen so far in this chapter are fairly crude. Many kinds of complications arise that make programs using them limited in scope and brittle. In this section we consider the source of the problems and explore the use of a Python module that solves some of them.

Problems with HTML pattern matching

Extracting the contents of header tags is one of the simplest things you might do with an HTML document. Even so, the expressions we used to accomplish that were not simple. Attempts to use regular expressions to selectively extract other parts of HTML pages will often run into more substantive difficulties.

One of the most irritating problems is that the people or programs producing the web pages from which you are extracting data may change the markup slightly after you've written your code. Where there was a space, for instance, there might be a "nonbreaking space" entity, written in HTML as . Some spaces may be eliminated, and line

breaks might be moved or deleted. Attributes in tags may also be added, removed, or changed. Every time this happens, you may have to update your program to accommodate the changes.

The real problem with HTML tags, though, is that they are nested. An 1i tag inside an ol tag can contain another ol tag that contains li tags, and so on. Likewise, there can be all sorts of span and div tags embedded inside other span and div tags. The documents are fundamentally tree-structured, like the example of the suffix tree in Chapter 4. Trees are recursive structures, so they can't be processed by linear methods.

In fact, browsers actually do convert the HTML they obtain from web servers into treestructured data, based on a standardized "document object model." In that model, each element has attributes, content, and children, just like the tags we've been describing. Because elements have children, the document object model is organized as a tree.

There are basic computer science techniques for handling recursive structures other than the tree pattern we encountered in Chapter 4. However, these can't handle HTML generally, for the same reasons the other Python libraries can't: many start tags in many HTML documents do not have corresponding end tags, and many complex parts of HTML files don't fit the simple pattern of tags with an opening, optional content, and a closing. Browsers must deal with these complexities, but it's too big a challenge for ordinary programming.

Structured HTML parsing: html.parser

Fortunately, Python's library contains a class that can deal with messy HTML content. Contained in the html.parser module, HTMLParser defines a framework that calls empty methods at significant points in its parsing of HTML text. The framework is fully implemented, though by itself it does nothing. All you have to do is define a subclass of HTMLParser that overrides some of its methods with definitions that implement the actions you want your application to perform.

The way to use your subclass is to create an instance of it (with no arguments), then provide it text with the feed(string) method. This method implements a form of buffering, so you can actually call it multiple times. When you are done feeding the instance, you call close on it. Methods called by an HTMLParser instance when it encounters the corresponding HTML construct include:

handle starttag(tag, attributes)

Called when <tag> is encountered, with tag the name of the tag (in lowercase) and attributes a list of name/value pairs

handle endtag(tag)

Called when </tag> is encountered, with tag the name of the tag (not including the slash)

handle startendtag(tag, attributes)

Called when a tag that is closed in its start tag is encountered (such as <hr/>r/>), with tag the name of the tag (in lowercase) and attributes a list of name/value pairs; if not overridden, it calls handle_starttag and then handle_endtag

handle data(data)

Called for text that is not a tag or something else special to HTML, with *data* a string containing that text

handle comment(data)

Called when a comment is encountered, with *data* the text of the comment

As a simple example, parsing <h1>a heading</h1> will result in the following calls:

```
handle_starttag('h1')
handle_data('a heading')
handle_endtag('h1')
```

There are a few other "handlers" that cover more obscure HTML items, which aren't mentioned here. Example 8-9 shows a subclass of HTMLParser that implements the outline generator in a more robust way than the one in Example 8-6.

Example 8-9. Using a subclass of html.parser.HTMLParser

```
class HTMLOutlineParser(html.parser.HTMLParser):
```

"""Show an indented outline of the headings in the HTML file provided as a command-line argument"""

```
# HTMLParser converts tag to lowercase
HeadingPat = re.compile('title|h([1-6])')
Indent = 4
def init (self):
    super().__init__()
    self.inheading = False
def handle starttag(self, tag, attrs):
    match = self.HeadingPat.match(tag)
    if match:
        self.inheading = True
        if match.group(1): # otherwise title & do nothing
             print('{0:{1}}[{2}]'.format(' ',
                                           self.Indent * int(match.group(1)),
                                           match.group(1)),
                   end=' ')
def handle data(self, data):
    if self.inheading:
        print(data, end=' ')
        # end=' 'so printing stays on this line in case
        # parts of the heading are nested inside other
        # markup, such as <i></i>
def handle endtag(self, tag):
    if self.inheading and self.HeadingPat.match(tag):
    # assuming heading tags (h1, h2, ...) cannot be nested
```

```
print()
                                # close tag
            self.inheading = False
if name == ' main ':
   if len(sys.argv) != 2:
       print('Usage: prognam file.html')
   else:
       filename = testfile
       parser = HTMLOutlineParser()
       with open(filename, encoding='utf8') as file:
            contents = file.read()
       print('File contains', len(contents), 'bytes')
        parser.feed(contents)
        parser.close()
```

XML

XML stands for the Extensible Markup Language. It defines some basic syntactic rules for tags, attributes, and content that are similar to, but stricter than, those in HTML. What it doesn't define are what tags and attributes a document can use. That's the extensible part: XML documents can contain any tags, attributes, and content. The tag names, their allowable attributes, what kinds of tags can be nested in what other kinds of tags, and other such details are specified by a schema written in a formal schema language, such as Document Type Definition[‡] and RELAX NG.§

Before looking at Python's facilities for processing XML files, we'll explore their structure in more detail. We'll also take a look at an XML datafile that we'll use in subsequent Python examples.

The Nature of XML

Roughly speaking, XML is the data equivalent of HTML. Its purpose is to support the text-based representation of structured data in well-defined formats. A great deal of data in bioinformatics, as well as in other fields, is available in XML form on the Web.

XML must adhere to very strict rules:

- The first line of an XML file is expected to specify the schema it uses.
- Tags with textual content must be completed by corresponding end tags.
- Attribute values must be quoted (single or double).
- A tag with no content may end with /> in place of a separate end tag.

§ See http://www.relaxng.org.

[‡] See http://www.w3.org/QA/Tips/Doctype and http://www.w3.org/QA/2002/04/valid-dtd-list.html.

The regularity enforced by these rules makes parsing XML files straightforward. That doesn't mean you'll want to write that code yourself, though, especially not when there are Python library modules to do it for you.

Other representation languages used in bioinformatics, such as the *Resource Description Framework* (RDF) and the related *Web Ontology Language* (OWL), are based on XML. Those languages are part of the more general Semantic Web initiative. Various parsers and even logical deduction mechanisms are available for them.

A variety of approaches to parsing XML files are possible. Some parsers process an entire file at once, while others parse incrementally. Some produce a tree structure that represents the XML content, while others generate "events." (The subclass of html.parser.HTMLParser defined in Example 8-9 implements simple event-based processing.) The event-based model is particularly useful when a large amount of XML text is to be searched for specific information. The tree-based model is appropriate where many actions will be performed on the results of parsing the entire file. Some modules can be used in several ways, and most have advanced features we won't cover here.

A parser understands a language's grammar, but it leaves it up to other code to process the syntactic elements it has extracted from its input. Python's library contains several event-based parsers, and XML's highly constrained syntax makes parsing an XML file straightforward. (Verifying that a schema's rules are followed in addition to the basic syntactic rules is a much more difficult challenge.) Parsing an XML file basically generates information about tag names, attribute name/value pairs, tag content, and the nesting relationships among the tags. It is up to the application to interpret what all that means.

Our main goal here is to demonstrate the nature of XML data and some of the ways it can be processed. We'll use two Python modules to parse some XML data downloaded from the Web. One, xml.etree.cElementTree, creates a tree for subsequent use. The other, xml.parsers.expat, is an interface to the event-based Expat C library.[‡] The

|| See http://www.w3.org/RDF.

#See http://www.w3.org/TR/owl-features, http://www.w3.org/TR/owl-guide, http://www.w3.org/TR/owl-ref, and (for the new features introduced by OWL 2) http://www.w3.org/TR/2009/WD-owl2-new-features -20090421.

^{*} See http://www.w3.org/2001/sw/.

[†] We've seen this kind of choice before, in processing FASTA files. We had the option of reading in all the sequences with their descriptions and then searching for sequences meeting certain criteria, or reading one sequence at a time and stopping after locating the one(s) meeting those criteria. The advantages and disadvantages here are the same. Processing an entire file means doing a lot of computation and producing a large data structure that the program can then use for repeated actions. An event-based scan of a file could potentially stop after a relatively small amount of processing and retrieve only a small amount of data; however, getting different data later would mean reparsing the file.

[‡] See http://www.libexpat.org.

cElementTree module will detect violations of the XML file's schema, but Expat will not—it is a nonvalidating parser. The xml package contains several other parsers, including some that work with the same document object model (DOM) that browsers use when interpreting HTML and executing JavaScript, but we won't cover those here.

An XML File for a Complete Genome

For this part's examples we are going to use an XML file representing an entire bacterial genome. The steps followed to obtain this are as follows (you can choose your own organism, of course; we're using a microbe genome because its XML file is of manageable size):

- 1. In a browser, go to http://www.ncbi.nlm.nih.gov/genomes/MICROBES/microbial taxtree.html.
- 2. Expand some taxa (in green) until you see individual organism names (in blue), e.g., the class Acidobacteria, about nine lines down.
- 3. Click on an organism name. For Acidobacteria only one organism name is shown: Acidobacteria capsulatum.
- 4. Click on the organism to get a page with an extensive summary of the information available about the organism and its genome.
- 5. Below the tabs visible in the upper-left corner, select "Protein Table" from the pulldown that is showing "Overview."
- 6. From the "Send to" pull-down, select "File."
- 7. Save the file wherever you want, perhaps naming it with the "UID" of the organism, as shown in the page title, and the extension .xml.

This will be a fairly large file. (The one for Acidobacteria capsulatum was 44 MB.) It has a rather deeply nested structure, which is summarized next. While you read the summary, it will help to refer to the display in the web page from which you downloaded the data to see how the two correspond.

The outline of the saved file is as follows. The numbers of lines listed are rough counts for the file Acidobacteria capsulatum, which identifies 3,377 proteins. The file consists of:

- The two lines identifying the XML version and the document schema that usually appear at the beginning of an XML file
- A few top-level start tags
- Some tags inside a Bioseq-set descr tag that provide basic information about the organism, its source, publication data, etc. (1,500 lines)
- Some tags inside a Bioseq-set seq-set tag that provide information about the genome (142,000 lines)

- A Seq-entry tag for each protein that describes that protein's sequence (600,000 lines)
- A Seq-annot tag for each protein that describes its annotations (155,000 lines)

The file uses a over 200 distinct tags and over 500,000 tags in total, as well as some attributes and tag content.



The following Python script counted the tags and distinct tags. It is very important to appreciate the power of the tools you are acquiring and learn to use them not just for what you consider "programming," but for small one-time tasks as well):

```
import re
pat = re.compile((r'</([A-Za-z-_]+)')
with open(filename) as file:
    closetags = pat.findall(file.read())
print('The number of total tags was {}, ' +
    'of which {} were distinct' .format(len(closetags), len(set(closetags)))</pre>
```

The ElementTree Module

The xml.etree.ElementTree module provides an easy way to read an entire XML document and create objects to represent its elements, attributes, and content. Its only disadvantage is the same as its strength: it reads the entire file. Whether to use an event-based parser or one that processes the entire file and creates a data structure from its contents depends on what your application will do with the parsed XML. If you had to process an extremely large XML document, or use the objects for many smaller ones together, reading their entire contents could cause problems. Searching for a particular element, for example, would suggest the use of an event-based parser.

For extreme situations or complicated processing patterns, one of Python's XML libraries that we aren't covering here might be more appropriate. There are also a few more advanced features of the ElementTree module that would allow partial processing to get something of a compromise between the two approaches.

Two Versions of the ElementTree Module

There are actually two versions of the ElementTree module: xml.etree.ElementTree and xml.etree.cElementTree. The first is written in Python, and the second is written in C with a Python interface. The C version's parser is quite a bit faster than the Python one, and the behavior of some of its methods differs in a few advantageous ways. In particular, its functions that return requested nodes return generators that produce them one at a time, whereas the Python versions return complete lists. For large numbers of nodes, a generator would be much more efficient. There being in this case no particular advantage to using the Python version or disadvantage to using the C version, we'll use the C one.

To avoid confusion between the module ElementTree and the class ElementTree. Element Tree, both in this discussion and the code examples, we'll import the module with the name Etree. In addition to being a shorter name, creating this alias for the actual module has the more significant benefit that the code will be able to work with either version of the module—we just have to change the import statement to import from whichever of xml.etree.ElementTree and xml.etree.cElementTree we want to use:

```
>>> import xml.etree.cElementTree as ETree
<module 'xml.etree.cElementTree' from</pre>
'/usr/local/lib/python3.1/xml/etree/cElementTree.py'>
```

Getting started with ElementTree

An instance of the class ETree. ElementTree represents a tree with features corresponding to the various aspects of XML. The tree consists of ETree. Element instances. Most of the ETree. Element Tree instance methods simply call the same method on its root Element—an example of delegation, as discussed in "Class decomposition" on page 186 (Chapter 5)—but a few provide additional functionality.

The module function ETree.parse(source) parses the XML text from source, a filename or a file open for reading, and returns an instance of ETree. Element Tree. To write a tree to a file, call its method write (destination), where destination is either a filename or a file open for writing.

ElementTrees are quite general and can be used for pretty much any kind of treestructured information. The module provides functions for building a tree programmatically, which you can use to convert hierarchical but non-XML data into XML, or to structure data your program is generating in preparation for its output as XML. You could also use it to construct trees for the program's own internal purposes that have nothing to do with XML. These are powerful capabilities, but we won't be discussing them any further. Our focus here is on manipulation of the tree created by the parsing functions.

We'll start by creating a tree for an XML genome file, such as one downloaded as described earlier in "An XML File for a Complete Genome" on page 302. We can see in Example 8-10 that ETree.parse returns an instance of ElementTree. The root of the tree is an Element. We can also see that while ElementTree instances don't give us anything more than the default information when the interpreter prints them, Element instances at least show the names of their tags.

```
Example 8-10. Basic ElementTree manipulations
```

```
>>> source = 'data/Entrez/sequences-24289.xml'
>>> tree = ETree.parse(source)
>>> tree
<ElementTree object at 0x18a3e70>
>>> root = tree.getroot()
```

```
>>> root
<Element 'Bioseq-set' at 0x18a7188>
```

The XML-related features of an Element are accessed as Python attributes, not with method calls. There are four:

tag

The name of the tag, as shown in the printout in Example 8-10

attrib

A dictionary of attribute names and values as extracted from the start tag

text

The text, if any, that appeared between the tag and its end tag

tail

The text appearing between the end tag and the start of the next tag, whatever that tag is

The Python documentation warns that although the current implementations use actual Python dictionaries for Element.attrib, that may not always be the case. It is recommended that the following Element methods be used instead of directly accessing attrib. They are defined to perform the same actions as the corresponding dictionary methods:

- items()
- keys()
- values()
- get(key[, default])
- set(key, value)

Navigating around an ElementTree

Pretty much everything a program does with a tree involves navigating around its nodes. Sometimes a program takes steps to get to a specific node. Sometimes it searches for one that meets specific criteria. Other times it *traverses* the entire tree, doing something to each node encountered (a Recursive Tree iteration, as described in Chapter 4). ElementTree and Element provide the following methods for these kinds of operations:

find(target)

Returns the first (immediate) child matching target

findall(target)

Returns a list of all the (immediate) children matching *target*, in the order in which they appear in the document

findtext(target[, default=None])

Returns the text of the first (immediate) child matching target, or default if no such element was found; an empty string is returned if a matching target is found that has no text content.

```
getiterator([tag=None])
```

Returns a generator that produces, from an element and all its descendants, all elements whose tag is tag; if tag=None or tag='*' the generator returns all of the elements

In addition, Element, but not ElementTree, provides the following method for navigating the tree recursively:

```
getchildren()
```

Returns a list of all the subelements of Element, in the order in which their tags appeared in the document

ElementTree, but not Element, also provides the getroot() method to get its root Element.

Example 8-11 shows a function based on the Filtered Count template from Chapter 4 ("Nested iterations" on page 126). The definition counts the number of nodes in an ElementTree subtree whose tag has a specified name.

Example 8-11. Counting the nodes for a specified tag

```
def count nodes(element, tagname):
    """Return the number of tagname nodes in the tree rooted at element"""
    for node in element.getiterator(tagnames):
        count += 1
    return count
```

Initial examination of the XML seemed to indicate that there was a series of Seq-entry tags, first one for the entire genome and then one for each protein. These contained core information. Later in the file there is a series of Seq-feat tags that follow in the file with sequence annotations. Let's see whether there are the same number of Seq-entry tags as Seq-feat tags:

```
>>> count nodes(tree, 'Seq-entry')
>>> count nodes(tree, 'Seq-feat')
10360
```

Apparently not. This is an easy way to interactively investigate simple hypotheses you might have about the structure of a deeply nested and/or large XML file after some preliminary examination using an editor. How about getting all tag names in the file, avoiding duplicates by using a set?

```
>>> tags = set()
>>> for element in tree.getiterator():
```

```
tags.add(element.tag)
>>> len(tags)
208
```

Let's count how many times each tag occurs:

Since there are 208 keys in this dictionary, we don't want to print the whole thing. We can first check the results obtained this way against the results obtained earlier using count nodes:

```
>>> counts['Seq-entry']
3379
>>> counts['Seq-feat']
10360
```

Now let's print out the counts of the 20 most common tags. This is another way to explore the nature of an XML file. There's no guarantee that a frequently occurring tag is only used inside a particular other tag—it could be used all over the file—but at least these counts are a hint. With some more work we could write a program that counted how many times each tag appeared within each other tag, and so on:

```
>>> sortedtags = sorted(counts.items(),
                         key=lambda item: item[1],
                         reverse=True)
>>> for item in sortedtags[:20]:
        print(item)
('Object-id', 27150)
('Seq-id', 23872)
('Object-id str', 20277)
('Seq-id_gi', 17116)
('User-field', 16898)
('User-field data', 16898)
('User-field label', 16898)
('Seq-loc', 13739)
('Segdesc', 13520)
('SeqFeatData', 10384)
('Seq-interval to', 10361)
('Seq-interval id', 10361)
('Seq-interval from', 10361)
('Seq-loc int', 10361)
('Seq-interval', 10361)
('Seq-feat_location', 10360)
('Seq-feat data', 10360)
('Seq-feat', 10360)
('Na-strand', 6984)
('Seq-interval strand', 6984)
```

Here are a few examples of method calls using the tree created in Example 8-10:

```
# all elements with protein names
>>> proteins = root.getiterator('Seqdesc_title')
>>> protein1 = next(proteins)
```

```
>>> protein1
                             # first protein is actually the whole genome
<Element 'Seqdesc title' at 0x1965218>
>>> protein1.text
'Acidobacterium capsulatum ATCC 51196, complete genome'
>>> next(proteins).text
'YkgG family protein [Acidobacterium capsulatum ATCC 51196]'
>>> next(proteins).text
'iron-sulfur cluster binding protein [Acidobacterium capsulatum ATCC 51196]'
>>> prot = next(proteins)
>>> print element(prot)
 Seqdesc title: cysteine-rich domain protein [Acidobacterium capsulatum ATCC 51196]
```

Example 8-12 shows a pair of function definitions used to print a subtree starting at a given Element. This is a straightforward use of the Recursive Tree Iteration template like the one in Example 4-26 (see "Recursive iterations" on page 128).

Example 8-12. Functions for printing subtree of an ElementTree

```
def print element(element, level=1):
    ""Print tag, content, and attributes of element, indented level
    spaces, but only if element has text or attributes"""
    # ignoring tail
    if element.text and not element.text.isspace():
        print(' '*level, element.tag, ': ', element.text.strip(), sep='')
    elif element.attrib:
        print(' '*level, element.tag)
    for attr in sorted(element.keys()):
        print(' '*(level+2), attr, '=', element.get(attr))
def print subtree(element, level = 0):
    ""Print the tags, attributes, and contents of element and all of its children,
    in depth-first order, starting with an indentation of level spaces"""
    print element(element, level)
    for subelt in element.getchildren():
        print subtree(subelt, level+1)
```

Example 8-13 shows what a printout using these functions would look like.

Example 8-13. Printing an ElementTree subtree

```
>>> descrs = root.getiterator('Segdesc source')
>>> print subtree(next(descrs))
Seqdesc source
BioSource
 BioSource genome: 1
    value = genomic
 BioSource org
  Org-ref
   Org-ref taxname: Acidobacterium capsulatum ATCC 51196
   Org-ref db
     Dbtag
      Dbtag db: taxon
      Dbtag tag
      Object-id
       Object-id id: 240015
```

```
Org-ref orgname
     OrgName
      OrgName name
       OrgName name binomial
        BinomialOrgName
         BinomialOrgName genus: Acidobacterium
         BinomialOrgName species: capsulatum
      OrgName mod
       OrgMod
        OrgMod subtype: 2
           value = strain
        OrgMod subname: ATCC 51196
       OrgMod
        OrgMod subtype: 35
           value = culture-collection
        OrgMod subname: ATCC:51196
       OrgMod
        OrgMod subtype: 255
           value = other
        OrgMod_subname: type strain of Acidobacterium capsulatum
      OrgName lineage: Bacteria; Acidobacteria; Acidobacteriales;
Acidobacteriaceae; Acidobacterium
      OrgName_gcode: 11
      OrgName div: BCT
```



Many other manipulations are possible using the small set of methods documented earlier. Experiment with them using a tree like this. While experimenting, it is best to assign names to results rather than just having them print in the interpreter—you never know how many thousands of lines will result from a method call! If you assign a name, you can check its length, or its first element, or some other feature before trying to print anything. You could also write functions that print only a limited number of items.

Example 8-14 illustrates a function that prints the information about a limited number of proteins starting at a specified number. This calls the print_subtree function of Example 8-12 and relies on the information that each protein's description's outermost tag is called Seq-entry.

Example 8-14. Printing information for a range of proteins

```
def describe_proteins(tree, limit=2, start=1):
    # start at 1 because 0 is the whole genome!
    iter = tree.getiterator('Seq-entry')
    # +1 to always skip entry for entire genome
    for n in range(start+1):
        next(iter)
    for k in range(limit+1):
        print('{:4}'.format(k+start))
        print_subtree(next(iter), 6)
```

Event-Based Processing

Before exploring one of Python's event-based XML parsers, we need to take an excursion to confront some unusual properties of event-based processing in general.

Function calls, exceptions, and the call stack

The usual pattern of function calling has the following characteristics:

- 1. When function A calls function B, it passes values for function B's parameters.
- 2. Function B returns a value to function A. (Either it executes all of its statements without encountering a return statement, in which case it returns None, or at some point it executes a return statement.)
- 3. When function B returns, function A continues from where it called function B.

Function calls are implemented by adding a data structure called a frame to the call stack for each function called. That frame contains (at least) the following critical information:

- Bindings for the function's local variables
- The address in the computer's memory of the machine-level instruction at which the function should continue when a function it has called returns to it
- The frame it should return to when it executes a return statement or finishes executing all of its statements

These stack frames implement the usual pattern of function call and return. When a function returns, its frame is "removed" from the stack. Actually, the only thing that really happens is that an index representing the current top of the stack is incremented so that it references the previous stack frame. If the function of the previous frame then calls another function, the new function's frame simply overwrites the frame of the one that just returned. Figure 8-1 illustrates the process.

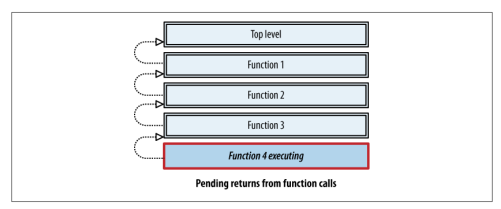


Figure 8-1. Function calling and stack frames

When an exception is raised, it returns control to somewhere further back in the stack. Unless the exception is caught within the same function in which it was raised, frames are removed from the stack, in a process called *unwinding*, until an except clause is encountered that matches the exception raised. This breaks the normal pattern of function call and return, since some of the "deeper" functions are simply abandoned without having a chance to execute the rest of their statements and return normally.

In the normal operation of the stack, each function returns control to the one that called it by making the caller's stack frame the current one. For control to return to the except clause of a try statement requires that there be another kind of entry in the stack to represent the clause apart from the stack frame of the function that contains the statement. When an exception is raised, Python unwinds the stack, checking every except clause entry along the way until it finds one that specified an exception class that matches the type of the exception being processed. Finally, it starts executing the statements in that except clause.

Almost everything about exception handling is the reverse of function calling:

- Each function call *adds* one frame to the stack, whereas raising an exception removes one or more frames from the stack.
- Function calls pass control to a *known* piece of code, whereas raising an exception tosses control up the stack with *no idea* of where it will end up.
- In the absence of an exception that bypasses the function call, a function *returns* control to where it was called from, whereas control is never returned to where the exception was raised.

In effect, raising an exception "calls" backward in the stack with a single argument to an unknown location, causing all the intervening stack frames to be discarded (see Figure 8-2). This is a strange beast indeed, yet one that plays critical roles. The lack of a direct connection between where the exception is raised and where it is caught, and the reverse direction of the frame movement in exception handling, is what makes it such a valuable tool.

The primary benefit of an exception handling mechanism is that it allows the separation of the code that detects an exceptional situation from the code that knows what to do about it. In particular, it allows library code to raise exceptions that are caught by application code that has called into the library. The library code that detects the exceptional situations will rarely know what actions it would be appropriate to take, while the code using the library will rarely be able to detect the exceptional situations.

Callbacks: Reversed function calls

There is another kind of reversed function call, known as a *callback*. It is not disruptive the way exceptions are: it is really just an ordinary function call used in a special way. Callbacks are a major feature of event-based processing where the processor, typically something from a library, calls a specific application function each time a corresponding

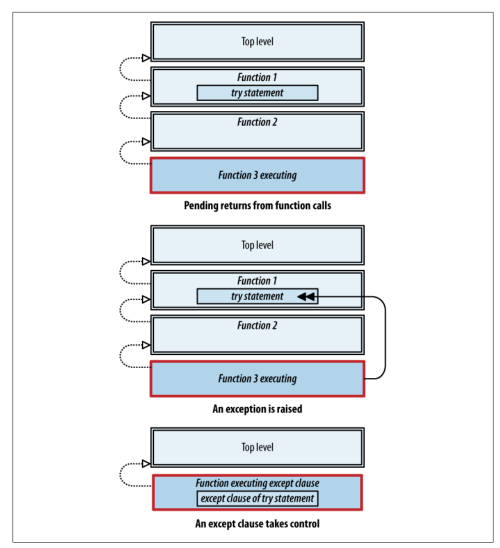


Figure 8-2. Stack frames and exceptions

event occurs during its processing. This places the processor in control of the computation while still allowing the application to define the actions to take at well-defined points.

Graphical user interface systems are the archetypal example. These systems monitor input devices such as mice and keyboards, constructing objects representing "events" for each keystroke and mouse click. It looks up each event in a table (a dictionary or something similar) to retrieve a description of the action to take in response to the event, usually expressed as a function to call. One of the challenges of this kind of architecture is giving application code a way to specify which function should be called when a particular icon is clicked, key is pressed, and so on.

Callbacks are the mechanism that allows the application to respond to processor events. The application defines functions and binds events to them. Then it turns control over to the event-based system. When an event occurs, the system "calls back" to the application via the function corresponding to the event. The technical definition of the system and its callback mechanism specifies what arguments will be passed to the function call through this mechanism. Some systems also allow applications to register additional values to be passed back to a function when the corresponding event occurs.

No new mechanisms are required to implement a callback system. Event-based processing simply reverses the roles of library and application code. The library provides functions for registering callback functions. After the application registers its callbacks with the event-based system, it passes control to it. Subsequently, the application functions play the kind of passive role that library functions usually do. The order in which they are called is undetermined since the calls are in response to input to the event-based system. Figure 8-3 illustrates this arrangement.

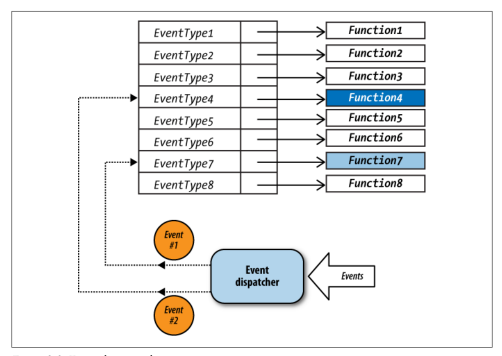


Figure 8-3. Event-driven architecture



Once you understand all of this, you'll see that the major difference in programming an application with an event-based system is the initial setup that associates functions with events. Some systems use assignment statements to do the binding, while others use function calls. Either way, the principle is the same.

Note that callbacks don't have to be functions named with a def statement—they can be "anonymous" functions created with a lambda expression. Just remember that lambda expressions cannot themselves include statements, though there's no problem with them calling functions that do. Like nested function definitions, lambda expressions have access to all the names in the scope in which they are defined. Therefore, they can pass information from the scope they are in as an argument to the function in another scope.

In a way, the html.parser.HTMLParser class we looked at earlier in this chapter implements event-based processing. Every time one of its instances encounters a tag in the HTML text it is parsing, it calls the method handle starttag. Every time it encounters a closing tag, it calls handle endtag. To implement an application using this class, you implement a subclass and override the class's "do-nothing" methods with definitions that perform your application's actions. This is a kind of event-based processing, with each kind of thing the parser recognizes causing the corresponding method to be invoked.

It feels as if html.parser.HTMLParser is in control, not your class, because it does all the parsing and event handling; your subclass just defines what to do with each event. In reality, though, the instance of your subclass does the parsing and event handling. It uses methods inherited from HTMLParser, including ones you may not even know about, but there is only one object doing the work. (In true event-based processing there is one object that generates events and calls event handlers and a separate object that handles the events.)

Programming for an event-based processor

Event-based architectures confront the programmer using them with two interesting challenges:

- How to stop the processing
- How to maintain application state between callbacks

These two problems stem from the same source. Typical applications are coherent in the sense that their conditionals, loops, iterations, and function calls constitute a seamless flow of computation. However, the application side of event-driven systems is fragmented: its functions are called by other code, at unpredictable times, and under unpredictable circumstances.

To set things up so an application exits in response to a certain event, you can simply have it call <code>sys.exit</code> from the callback function. However, it is sometimes the case that the application wants to take back control from the event-based system. Doing so requires returning past where the event loop was invoked, back into the application code. We can do this by raising an exception in a callback and catching it where the event system was invoked:

```
def stop(args):
    raise StopIteration

# set up callbacks, etc.
try:
    call_external_event_loop()
except StopIteration:
    pass
# do more
```

The code is straightforward, except that its try statement looks misleadingly like a typical one designed to handle errors. StopIteration was chosen as the exception class instead of the more general Exception to provide at least a hint of the purpose of the try. The strange function stop, which simply throws the exception, would be registered as the callback for an event. Unless something in the event processor happens to catch StopIteration, as soon as stop is invoked from the event loop it will throw the exception. The stack will then be unwound all the way through the event loop and back to the try statement that invoked it—awkward, but effective.

The second problem requires a way for separate functions to share state. Normally, names are bound to values through function calls and assignment statements. The names in one function are independent of the names in any other function, even if they are spelled the same. Likewise, names assigned inside a function are independent of names assigned at the top level of a module that contains it. Event-based architectures present a problem for this kind of arrangement because the application functions don't call each other, but instead are called by the external system. They are therefore denied the function call arguments mechanism by which they normally pass information to each other.

We addressed this problem earlier by using a class to encapsulate all the necessary state together with the methods that manipulate it. Many methods of the GenBank parser class we created in Chapter 5 (see Example 5-4 and the surrounding discussion) used the same variables. Instead of cluttering up the code with many parameters in method definitions and arguments in method calls, all the definitions just accessed the fields of the instance. A method of a subclass of html.parser.HTMLParser may need values computed in a previous call or in a call to a different event-handling method. Storing values in the fields of an instance makes them available to all the methods of the instance's class, whatever happens between calls to them.

GenBankParser was an ordinary class that defined all its own methods. The HTMLOutlineParser class that we defined in Example 8-9 is a subclass of the class that

generates the events. In both cases, all calls stay within the context of a single instance. When an external event-based processor is used, we need a way to tie the event handlers to instance state. Python gives us two easy ways to do just that:

- Define one function inside another (usually with a lambda expression, but one def statement can appear inside another). The inner function, when executed, has full access to the local state of the function in which it was defined. If the outer function is a method, the inner function can access the instance through the name self, just as the outer method can.
- *Use a bound method.* You have undoubtedly seen interpreter output of the form:

```
<bound method ClassName.method name of object>
```

This is what you get when you type

```
>>> name.method name
```

with no parentheses, where name is the name of an instance of ClassName and method name is the name of a method of ClassName. The way this works might surprise you at first, but it is just binding at work as usual. Because method name is accessed through the value of an instance, its value is the value that *name* has within the instance's namespace, with self bound to that instance. In other words, a bound method carries around with it the instance to which it is bound.

Example 8-15 presents a tiny class that demonstrates how these two approaches work.

Example 8-15. Access to outer scope from inner functions

```
"""Hold a value, demonstrating some aspects of nested scope"""
    def __init__(self, v=None):
        self.val = v
    def get(self):
        return self.val
    def set(self, v):
        self.val = v
    def setter(self):
         """Return a function that can be called to change the value held by the instance"""
        return lambda newval: self.set(newval)
    def str (self):
        return str(self.get())
    def __repr__(self):
    return 'value({})'.format(self.get())
>>> v = Value()
>>> v
value(None)
>>> v.set(0)
```

```
>>> v
value(0)
>>> fn = v.setter()  # with parentheses
>>> fn(9)
>>> v
value(9)
```

The function returned by Value.setter is an ordinary function with one parameter, not a method. When it is called, it calls self.set with its parameter as the argument. The point of this is that every function (and method) has access to all the names available where it is defined. If it is defined inside another function (or method), it has access to that other function's parameters and to any names the other function has assigned. This demonstrates the first of the previously mentioned techniques. Using the same Value class, here's a demonstration of the second technique:

```
>>> fn = v.set  # no parentheses
>>> fn
<bound method value.set of value(9)>
>>> fn(4)
>>> v
value(4)
```

expat

The xml.parsers.expat module is a straightforward incremental event-based parser based on the widely used Expat Clibrary. To use it, you define a few callback functions, create an instance of a parser object, assign some of its fields to your callback functions, and feed the parser text. The text can be fed all at once or in segments. To create a new instance, use the following method:

```
xml.parsers.expat.ParserCreate([encoding])
```

Creates an instance of xml.parsers.expat.xmlparser for text with the encoding specified by *encoding*. *encoding* defaults to None, with the only other options being 'UTF-8', 'UTF-16', 'ISO-8859-1' (Latin1), and 'ASCII'.

To parse XML text, use one of the following two methods with the instance returned from ParserCreate:

```
Parse(string[, endflag])
```

Parses *string*, which may be empty; *endflag* (default False) must be True on the last call to this method

ParseFile(readable)

Parses the contents of *readable*, which can be any object that provides the method read(*nbytes*); since it reads bytes, if *readable* is an opened file, it must have been opened in binary mode

§ See http://www.libexpat.org.

The most common event handlers are similar to the event-handling methods of html.parser.HTMLParser, because HTML is similar to (though much less wellstructured than) XML. However, they are fields, not methods: you assign them rather than overriding them in a subclass. Their values are functions.



Remember, functions (including lambda expressions) are objects, objects are values, values can be named, and fields of an instance are one kind of name. A function can therefore be assigned as the value of a field

The three event handler fields that will serve most needs are as follows, with parser an instance of xml.parsers.expat.xmlparser:

StartElementHandler

The function to be called when a start tag is encountered, with its first argument the name of the tag and its second a dictionary of attribute names and values

StopElementHandler

The function to be called when an end tag is encountered, with the name of the tag as its sole argument

CharacterDataHandler

The function to be called for text encountered outside of a start or end tag, with the text as its sole argument

Other than callbacks, the instance has a number of fields that control its behavior, most of which are fairly technical. This one is worth knowing about:

buffer text

Controls whether text parsing should stop at every newline (if False, the default), calling CharacterDataHander, or should join consecutive text lines together and pass them as one large string to CharacterDataHander; this can be set at any time, even after some text has been parsed, and can make parsing considerably more efficient

Obtaining a single piece of information

We'll start with something simple. We downloaded this enormous file, but we want to extract the content of just one tag from it. Example 8-16 shows how we would extract the value of the first (and, as it turns out, the only) occurrence of the tag <Seqdesc title>. It uses the exception handling trick discussed earlier to stop the program. The program is simple because it doesn't need to keep state between callbacks, and all it does when it receives the StopIteration exception is print the information that was included in the exception when it was created. This assigns event handlers to lambda expressions inside its function definitions: while the parameters of a particular kind of handler are predefined, the lambda expression doesn't have to use them all, and it can call other functions with any combination of argument values.

```
import xml.parsers.expat
def handle start(p, target, name):
    """Install data handler when target tag is encountered"""
    if name == target:
        # now install handler for data tag
        p.CharacterDataHandler = lambda data: return data(data)
def return data(data):
    """Stop searching, since the target has been found;
     installed when the target start tag has been encountered"""
    raise StopIteration(data)
def lookup(filename, target):
    p = xml.parsers.expat.ParserCreate()
    # install handler for start tags
    p.StartElementHandler = \
        lambda name, attrs: handle start(p, target, name)
        # one way to "forward" a parameter to an outside function
    with open(filename) as file:
        try:
             while True:
                 p.Parse(file.read(2000))
                                                 # read 2000 bytes at a time
        except StopIteration as stop:
                                                 # expected!
             return str(stop)
```

Extracting a few pieces of data

Next, we'll extract the genus and species names of the organism whose genome is in the file. The XML for this looks like:

Example 8-17 shows an implementation similar to that in the previous example. What's different about this one is that we can't just stop when we find one tag—we need to find a second one too (first the genus, then the species). Because we need to maintain state between callbacks, we must define a class and assign the callback fields to methods of one of its instances. The code starts by installing the typical three expat parser callbacks. This demonstrates the way expat is typically used.

```
Example 8-17. Getting the content of several XML tags
```

```
import xml.parsers.expat
class GenomeBinomialParser:
    ChunkSize = 4000
                                       # number of bytes to read at a time
    def init (self):
        self.parser = xml.parsers.expat.ParserCreate()
        self.parser.pbuffer text = True
        # don't break text content at line breaks
        # Install the three standard callbacks
        self.parser.pStartElementHandler = self.start element
        self.parser.pEndElementHandler = self.end element
        self.parser.pCharacterDataHandler = self.char text
    def start_element(self, name, attrs):
        """Record name of start tag just encountered"""
        self.current tag = name
                                      # ignore attrs
    def end element(self, name):
        """Stop parsing when species name has been encountered"""
        if name == 'BinomialOrgName species':
            raise StopIteration
    def char text(self, text):
        """If current tag is for genus or species, record its content"""
        if not text.isspace():
        # checking for whitespace because this function gets called
        # for both tag content and whitespace between the end of
        # one tag and the beginning of the next
            if self.current tag == 'BinomialOrgName genus':
                 self.genus = text.strip()
            elif self.current tag == 'BinomialOrgName species':
                 self.species = text.strip()
    def find binomial(self, filename):
        """Return (genus, species) as found in the XML genome file named filename"""
        # read only ChunkSize characters at a time in case we find it relatively early
        # and to not take up an enormous amount of memory with file contents
        with open(filename) as file:
            trv:
                 while True:
                     self.parser.Parse(file.read(self.ChunkSize))
            except StopIteration:
        return self.genus, self.species
if __name__ == '__main__':
    p = GenomeBinomialParser()
    print(p.find binomial(sys.argv[1] if len(sys.argv) > 1
                            else 'data/XML/Example.xml'))
```

Getting the content of all tags with a specified name

In the XML file we are parsing, the names of the proteins are inside tags named Seqdesc_title. Example 8-18 shows a class that extracts a list of the names of all the proteins in the file.

Example 8-18. Getting content from all occurrences of a tag

```
import xml.parsers.expat
class GenomeProteinNameParser:
    def init (self):
        self.titleflag = False
        self.proteins = []
                                                    # a form of collect iteration
        self.count = 0
                                                    # just for user feedback
        self.parser = xml.parsers.expat.ParserCreate()
        self.parser.buffer text = True
        # don't break text content at line breaks
        self.parser.StartElementHandler = self.start element
        self.parser.EndElementHandler = self.end_element
        self.parser.CharacterDataHandler = self.char data
    def start element(self, name, attrs):
        """Keep a count of all Segdesc_title tags and print a period to the terminal
        every 250 tags; if name == 'Seqdesc_title', turn on titleflag & ignore attrs"""
        if name is Seqdesc title, ignoring attrs""
        if name == 'Seqdesc title':
            self.titleflag = True
            self.count += 1
            if not self.count % 250:
                 print(self.count, file=sys.stderr)
                 # sys.stderr so not buffered
    def end element(self, name):
        """Set titleflag to False as soon as any end element is encountered"""
        self.titleflag = False
                                                    # regardless of close tag
    def char data(self, text):
        """If last start tag was a Segdesc_title, then add text to the list of protein names"""
        if self.titleflag:
                                                    # turned on by start_element
            self.proteins.append(text.strip())
    def get protein names from file(self, filename):
        print('Parsing...')
        with open(filename, 'rb') as file:
                                                    # ParseFile requires bytes
            self.parser.ParseFile(file)
                                                   # parse entire file
        return self.proteins
def print protein names(filename, number):
    p = GenomeProteinNameParser()
    protein names = p.get protein names from file(filename)
print('\n\nFound {} proteins; the first {} are:\n'.
```

```
format(len(protein names), number))
   for protein name in protein names[:10]:
       print(protein name)
if name == ' main ':
   _____default filename = 'data/XML/AcidobacteriumCapsulatum.xml'
   print protein names(sys.argv[1] if len(sys.argv) > 1
                       else default filename,
                        sys.argv[2] if len(sys.argv) > 2 else 10)
The output from running this program with no arguments is:
    Parsing...
    250 ...
    500 ...
    750 ...
    1000 ...
    1250 ...
    1500 ...
    1750 ...
    2000 ...
    2250 ...
    2500 ...
    2750 ...
    3000 ...
    3250 ...
    Found 3383 proteins; the first 10 are:
    Acidobacterium capsulatum ATCC 51196, complete genome.
    cellulose synthase operon protein YhjQ [Acidobacterium capsulatum ATCC 51196]
    hypothetical protein ACP 0957 [Acidobacterium capsulatum ATCC 51196]
    hypothetical protein ACP 0614 [Acidobacterium capsulatum ATCC 51196]
    hypothetical protein ACP 0540 [Acidobacterium capsulatum ATCC 51196]
    putative membrane protein [Acidobacterium capsulatum ATCC 51196]
    hypothetical protein ACP 0928 [Acidobacterium capsulatum ATCC 51196]
    type I restriction-modification system, S subunit [Acidobacterium capsulatum
    ATCC 51196]
    putative flp pilus assembly protein CpaB [Acidobacterium capsulatum ATCC 51196]
    glycosyl hydrolase, family 3 [Acidobacterium capsulatum ATCC 51196]
```

Tips, Traps, and Tracebacks

Tips

- You don't have to fit everything you are trying to do with HTML tags into a single regular expression. You can use several, or you can use one or more together with ordinary str methods.
- While developing a program, add functions for each step you recognize, even if you don't see anything for that step to do just yet. You can just use pass as the entire body of the definition, or just have it return its argument. Later, as you extend

- the program, you will often find that the structure is already there and you'll just have to expand the definitions of one or more existing functions.
- Once you've got a program working with a set of web pages you want to process, make sure to try it on some similar web pages. Those experiments will often reveal cases you didn't handle in your regular expressions or function definitions, or bugs present in your program that weren't hit by the pages you were using initially.

Traps

 Don't just print the results of accessing the children of an ElementTree element there could be thousands. First, assign a name to the result and check how many there are.

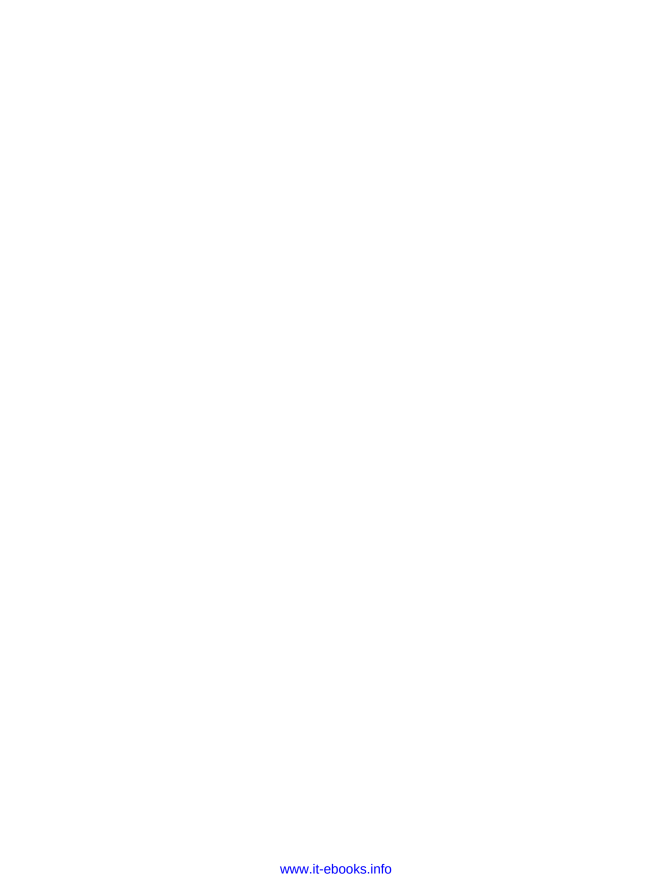
Tracebacks

Here are a few representative error messages:

UnicodeDecodeError: 'ascii' codec can't decode byte 0xc2 in position 1784: ordinal not in range(128)

An attempt was made to read a Unicode character from a file opened with the default ASCII encoding rather than encoding='utf-8'.

xml.parsers.expat.ExpatError: not well-formed (invalid token): line 1, column 1 Invalid XML appears at the beginning of a file; perhaps a blank line.



Web Programming

This chapter introduces a variety of Python facilities for manipulating URLs, opening documents in web browsers, submitting HTTP requests to web servers, and executing programs on web servers to respond to HTTP requests. Along the way we will also look at the foundations of the socket technology that underlies web interactions and the basics of constructing HTML forms to use as an interface to server-based programs.

Manipulating URLs: urllib.parse

The urllib.parse module provides functions for manipulating URL strings. The general form of a URL is:

scheme://network location/path;parameters?query#fragment

The fragment, which doesn't usually appear if the URL includes a query, is a reference to a particular place within a web page. You may not have seen or noticed the parameters portion of a URL before; it is not usually part of URLs visible to a browser's user, appearing—relatively rarely—as part of the value of href attributes in HTML tags. The network_location can be further dissected into the following components:

username:password@hostname:port

The *username/password* combination is a way of supplying login information to sites that accept this primitive kind of authentication. The *port* needs to be included whenever the server program that responds to a request for the given *scheme* is listening on a nonstandard port number. We'll see plenty of concrete examples of various forms of URLs in this chapter.

Certain characters are "reserved" for use in URL syntax and may not be part of the *query* component. These characters are represented by a percent sign (%) followed by their hexadecimal equivalent. You have undoubtedly noticed these in URLs, though you may not have realized what they were. Table 9-1 lists the characters and their URL equivalents. This translation is generally called *URL encoding*, but the official web standards call it *percent-encoding*.

Table 9-1. Reserved characters and their URL-encoded equivalents

!	*	"	'	()	;	:	@	&
%21	%2A	%22	%27	%28	%29	%3B	%3A	%40	%26
=	+	\$,	/	?	%	#	[]
%3D	%2B	%24	%2C	%2F	%3F	%25	%23	%5B	%5D

In addition, for various reasons, spaces in are not allowed in URLs. Sometimes they are translated to plus signs (+) and sometimes to their hexadecimal equivalent, %20. Note that URL encoding is *not* the same as encoding of HTML entities such as & or, equivalently, &. Not only is the context completely different, but, to confuse things further, URL encoding uses hexadecimal numbers while HTML entities are encoded in decimal.

Disassembling URLs

Extracting a URL's components is a miniature parsing problem with enough complexity that Python provides library functions that implement it. You will want to use these instead of writing your own code:

```
urllib.parse.urlparse(string[, default scheme[, fragmentflg]])
```

Returns an instance of urllib.parse.ParseResult, which looks like a six-element tuple whose elements correspond to the parts of the URL named in the preceding section. URL components not present in *string* are represented by empty strings, but if there is no scheme in string the tuple will contain default scheme (if specified). If fragmentflg is false, a fragment in string is ignored.

Punctuation used to delimit fields, such as the :// after the scheme name and the? before a query, are not included in the results. However, the initial slash of a path (if present) is retained, because in URLs that represent paths—those using a file scheme, in particular—an initial slash indicates an absolute path as opposed to a relative one (more on this later). In addition to its functioning as a six-element tuple the instance also has the read-only attributes shown in Table 9-2, the first six of which are shown in the order in which they appear in the tuple.

Table 9-2. Attributes of a ParseResult

Attribute	Description
scheme	URL scheme specifier
netloc	Network location
path	Hierarchical path
params	Parameters for last path element (uncommon)
query	Query component

Attribute	Description
fragment	Fragment identifier
username	Login information
password	Login information
hostname	The actual host without the login and port
port	Port number (integer) if present in location

urllib.parse.unquote(string)

Returns a copy of string with all %xx sequences replaced by their one-character equivalents.

urllib.parse.unquote plus(string)

Just like urllib.parse.unquote, except that it replaces plus signs with spaces.

Assembling URLs

Programs that work with web technologies frequently construct URLs from details specified as parameters of programs and functions or fields of classes and instances. The urllib.parse module provides a full range of functions for constructing URLs:

urllib.parse.urljoin(base, url[, allow_fragments])

Returns an absolute URL string constructed by adding to url any parts it is missing from the corresponding part of *base* (if present)

urllib.parse.urlunparse(parts)

Returns a URL string constructed from parts, which is typically a ParseResult but could be any six-item iterable

urllib.parse.urlencode(data)

Converts data, which is either a dictionary or a sequence of two-element tuples, to a URL-encoded string ready to be used as the argument portion of a query; the result is a string composed of key=value pairs separated by ampersands, with both key and value quoted by calling urllib.parse.quote plus

urllib.parse.quote(string)

Returns *string* with any special characters replaced by the corresponding %xx form to allow those characters to be included in a URL, especially in a query

urllib.parse.quote plus(string)

Just like urllib.parse.quote, except that it replaces spaces with plus signs instead of %20

urllib.parse.urldefrag(url)

Returns a pair containing a copy of url without its fragment as its first element and its fragment as its second; the second element will be an empty string if url does not contain a fragment

Opening Web Pages: webbrowser

The webbrowser module provides simple mechanisms for displaying documents or query results in a web browser. The module's functions and methods all take a URL as their first argument and instruct the user's browser to display it.



The functions in webbrowser give Python applications an easy way to use a browser as a presentation mechanism. An application can write some results into a text or HTML file to be displayed in the browser, submit a query through a browser, or just open a page of HTML-formatted documentation. Many other uses are also possible—all the program needs to do is provide a URL.

Module Functions

Following are descriptions of the webbrowser module's functions. All browser actions are subject to the browser in question's ability to perform those actions in the specified way with the user's current settings. For instance, if a function is described as opening a new tab, the actual action might be opening a new window. The functions are:

```
webbrowser.open(url[, new=0[, autoraise=1]])
```

Displays url in a browser window (the current browser window if new is 0, a new browser window if it's 1, and a new browser tab if it's 2); the window will be raised unless autoraise is false

```
webbrowser.open new(url)
```

Opens url in a new browser window (otherwise, it's opened in the current browser window)

```
webbrowser.open new tab(url)
```

Opens *ur1* in a new browser tab (otherwise, it's opened in the current browser tab)

For example, to open the *python.org* home page:

```
webbrowser.open('http://www.python.org')
```

A function that opens the documentation for one of the core library modules follows:

```
def browse module doc(module name):
   webbrowser.open('http://docs.python.org/py3k/library/' +
                    module name + '.html')
```

Alternatively, you could replace 'http://docs.python.org' with 'file://' and the full path to where the documentation is in your Python installation (beginning with a slash, /, to make it absolute).

On some platforms, opening a URL may open it in some other kind of application. The type of URL—for example, ftp vs. http—may affect this behavior. If the URL is a path to a file (whether or not it begins with *file://*), the system might choose an application based on the file's extension or one associated with that specific file. For instance, you

may have configured your system to open all .py files in IDLE, in which case webbrowser.open('/full/file/path/name.py') may open name.py in IDLE. This makes the webbrowser module useful for many scenarios beyond those involving browsers.*

Constructing and Submitting Queries

An application can use webbrowser.open to submit queries like the ones you see in your browser's address bar when you submit a query through a website. The only difference between a query and a URL for an HTML document is that the query is a path to a program on the client that gets its arguments from the part of the URL after a question mark. For example, there's a package index at http://pypi.python.org/pypi that lists information about a large assortment of externally available Python software. If you were to type something into the search box at the top of the page and submit the query, the URL the browser would send to the *python.org* server would be of this form:

```
http://pypi.python.org/pypi?%3Aaction=search&term=text&submit=search
```

where text is whatever you typed in and %3A is the URL encoding for a semicolon. If your entry has more than one word, the individual words will be separated by plus signs. (In other words, the spaces in the multi-word query are replaced by plus signs, as described earlier.)

Example 9-1 shows a complete program for running a search of the Python Package Index from the command line.

Example 9-1. Searching the Python Package Index

```
"""Search the Python Package Index for one or more words"""
import sys
import webbrowser
import urllib.parse
urlpart1 = 'http://pypi.python.org/pypi?;action=search&term='
urlpart2 = '&submit=search'
def search pypi(words):
    webbrowser.open(urlpart1 + urllib.parse.quote plus(words) + urlpart2)
if name == " main ":
    if len(sys.argv) < 2:</pre>
        print("Usage: pypi term ...")
    else:
        search pypi(sys.argv[1:])
```

^{*} Technically, using webbrowser for file:// URLs is unsupported and nonportable behavior, but if it does what you want, you might as well use it. Just make sure to note in your files that the behavior is undocumented and might well change in the future.

Constructing and Viewing an HTML Page

There are many ways to construct HTML pages. For instance, you could create an HTML document that includes & placeholders and use string. Template (see "String Utilities: string" on page 240 in Chapter 6) to replace them. (This is an ideal use for string. Template .) Another approach is to use format and assign some formatting strings for use in functions.

Example 9-2 shows an extension to the program for extracting links from HTML (shown in "Searching HTML text" on page 290). Assume you have run that program to produce some files that contain the pairs of URLs and text the program outputs. What this example does is use some of those files to produce a simple HTML file by putting each filename in an h3 tag, making a new a tag for each URL, and enclosing the a tags in a numbered list. (There's a good chance that some of the links picked up by extract links won't be meaningful or useful. In that case, you can always hand-edit the resulting file, or if there is a pattern to which links are useful and which aren't, you can filter them out either in that program's get all tags function or in this one's make html for file.)

Example 9-2. Making a web page with links extracted from HTML files

```
import sys
import webbrowser
html start = ('''<!DOCTYPE html PUBLIC</pre>
          "-//W3C//DTD XHTML 1.0 Transitional//EN"
          "http://www.w3.org/TR/xhtml1/DTD/xhtml1-transitional.dtd">
<html>
<head>
<title>Collected Links</title>
</head>
<body>
''')
html end = '''
</body>
</html>
filetagformat = \n<h3>{0}</h3>\n'
atagformat = '<a href="{0}">{1}</a>\n'
def make html file from files(filenames, outputfilename='links.html'):
    with open(outputfilename, 'w') as outfile:
        outfile.write(html start)
        for filename in filenames:
            make html for filename(filename, outfile)
        outfile.write(html end)
    return outputfilename
def make html for file(filename, outfile):
    outfile.write(filetagformat.format(filename))
```

```
outfile.write('\n')
with open(filename) as infile:
    for tag in get_all_atags(infile.read()):
        if tag[1]:
            make_html_for_tag(tag, outfile)
outfile.write('\n')

def make_html_for_tag(tag, outfile):
    outfile.write(atagformat.format(tag[0], tag[1]))

if __name__ == "__main__":
    if len(sys.argv) < 2:
        print("Usage: browse_links filename ...")
    else:
        webbrowser.open(
            'file://' +
            os.path.abspath(make_html_file_from_files(sys.argv[:1]))
            '</pre>
```

Web Clients

The webbrowser module we looked at earlier provides a simple way to request the opening of a URL. Its functions return shortly after they are called, leaving a browser or some other application outside of Python to fulfill the request. In contrast, the urllib.request module provides a variety of sophisticated services for interacting with web servers from within Python itself.

The most generally useful functions in the module are:

```
urllib.request.urlopen(url[, data[, timeout])
```

Performs the request specified by *url*, giving up after *timeout* seconds if specified; if a *data* argument is provided, it should be in the form returned by *urllib.parse.urlencode*, and it is supplied as the contents of a POST request (otherwise, the *url* itself would include the request's parameters); returns an object with the functionality of a file object and two additional methods:

```
urllib.request.geturl()
```

Provides the URL of the actual resource retrieved; possibly redirected from the original request

```
urllib.request.info()
```

Returns an instance of http.client.HTTPMessage, with information about headers, etc.

```
urllib.request.urlretrieve(url[, filename[, reportfn[, data)
```

Performs the request specified by *url*, returning a two-element tuple containing the path to the file where the contents of the response was saved and the headers that would be returned by *urllib.request.urlopen(url).urllib.request.info()*; optional arguments are as follows:

filename

The path to the file in which to save the guery response; otherwise, a temporary file is used

reportfn

If present, will be called when the network connection is established; then for each "block" of data read from the response, with three arguments: the number of blocks transferred so far, the number of bytes in a block, and the total size of the file

data

If present, will be used as the contents of a POST request (and should be constructed by calling url.parse.encode); otherwise, query arguments must be provided as part of the URL itself



Be careful using a string returned from urllib.request.urlopen, especially with regular expressions—the file-like behavior of the object returned provides a stream of bytes, not Unicode characters. Any read operation from the stream will therefore return a bytes, not a str. The simplest thing to do is just call str on the input, which will work if the bytes represent characters in the default encoding. You can specify an encoding by either giving it as the second argument to str or calling the decode method on the bytes, with the encoding as its argument.

If you try to use a string for a regular expression to match a bytes target, you'll get this error:

TypeError: can't use a string pattern on a bytes-like object

You could also use a bytes instead of a raw string for the regular expression, as long as the rest of the code is changed wherever necessary. The main point is that what go in and out over a network are bytes.

Making the URLs in a Response Absolute

URLs in a web page do not necessarily have a scheme (e.g., http://) or a hostname (e.g., www.python.org), making them pretty much useless as links. A URL without a scheme and host is known as a relative URL, because its actual address is derived from the site and path at which the document containing the link was found. If a program such as that shown in Example 9-2 is supposed to do something useful with these links, they must first be converted to full URLs so that they can be used outside of the context of the page on which they were found.

A relative address that begins with a slash (/) is treated as if it were prefixed with the same protocol and site as the page on which it is found. This is similar to how paths in filesystems are interpreted: an initial slash denotes the "root" directory, and any address that does not begin with a slash is considered relative to the directory that contains its document. If the address is just a filename, such as other.html, clicking on the link will look for that file in the directory in which the current document was found.

If there are any slashes inside the address, everything through the last slash is treated as a path starting with a subdirectory of the one containing the link's document. For example, a link *images/small/img1.jpg* in a web page at this URL:

http://www.example.com/top/middle/demo.html

would refer to a file at this location:

http://www.example.com/top/middle/images/small/img1.jpg

In contrast, because it begins with a slash, the link /images/small/img1.jpg would instead refer to a file at this location:

http://www.example.com/images/small/img1.jpg

Expanding relative URLs to absolute ones could require a lot of code to cover all the possible permutations of which parts are present in a URL and which aren't. This would get complicated quickly. There's no need to bother with direct string manipulation, though, because the urllib.parse module described at the beginning of this chapter gives us the tools we need to disassemble and assemble URLs. The function urllib.parse.urljoin is all we need here. It's not even necessary to test whether a link's address begins with a scheme and a host before calling urllib.parse.urljoin—the function works with whatever parts of a URL are present in the first argument. Any parts present in both arguments get their values from the second argument.

One problem with the programs we have been using up to this point is that they process saved HTML files. Although they know the names of the files, they have no information about the original source of their content. What we'll do now instead of saving and then reading HTML files is use urllib.request.urlopen to read web pages directly. We won't even bother writing them to files. Then, when the pages are read, the code will know their sites and paths and be able to use them to adjust the links they contain. The command-line arguments will be URLs from which to read—HTML pages or requests. (A long list of URLs can be provided by command-line input file redirection.)



Since web page contents are returned as bytes objects, not strings, if you want to write the modified contents to a file you'll have to either first convert them to strings or open the file in 'wb' mode.

Constructing an HTML Page of Extracted Links

Let's add some code to Example 9-2 to make the URLs in the document absolute. That way, the files the program constructs will be usable on their own. Example 9-3 shows the functions that replace those in Example 9-2. Everything else stays the same, except that the __main__ code calls make_html_file_from_urls instead of make_html_file_from_files. Also, the program must import urllib.parse and urllib.request.

Example 9-3. Constructing a web page directly from requests

```
import urllib.parse
import urllib.request
def make html file from urls(urls, outputfilename='links.html'):
   with open(outputfilename, 'w') as outfile:
        outfile.write(html start)
        for url in urls:
           make html for url(url, outfile)
        outfile.write(html end)
   return outputfilename
def make html for url(url, outfile):
   outfile.write(filetagformat.format(url))
   outfile.write('\n')
   for tag in get all atags(str(urllib.request.urlopen(url).read().decode())):
        if tag[1]:
           make html for tag(url, tag, outfile)
   outfile.write('\n')
def make html for tag(url, tag, outfile):
   outfile.write(atagformat.format(make absolute(url, tag[0]), tag[1]))
def make absolute(url, tagaddress):
   return urllib.parse.urljoin(url, tagaddress)
```

All the links in the HTML file produced by this revised program are absolute. You can open the resulting file in a browser and click on the links to go to their pages or submit their requests.

Downloading a Web Page's Linked Files

Occasionally, you may come across a web page that's full of links to files you would like to download. Some browsers, browser extensions, and applications provide tools for doing this. However, you might want to do this in a program to customize exactly which links are downloaded and where they get stored, or to more easily run the program as part of some automated task. It's easy to modify the code from the previous examples in this chapter for this purpose. We already know what the necessary steps are:

- 1. Parse command-line options.
- 2. Fetch a document or submit request.
- 3. Read from the request.
- 4. Convert the bytes read to a string.
- 5. Extract all links.
- 6. Save documents to a file.

There is one import twist we'll add to this program, though: we don't want to download *all* the links from a page, just those with certain extensions. We have no idea what kinds of documents, requests, media, and just plain useless things might appear on a web page, but we typically know what kinds of links we do want to download: perhaps all *.zip* and *.pdf* files, or all *.html* links. With that in mind, we'll set things up so that the command-line arguments for the program are a URL and one or more extensions to extract.

The program will not handle URLs with queries that are contained in the results returned by urllib.request.urlopen. Query URLs usually don't indicate the kinds of files they will obtain, and they usually return new HTML pages with their results rather than something you would want to save as a file. (Well, you might want to save them, but we've already handled various aspects of that, and you can always click on a request URL, save the file, and run this program, giving it a file URL with its path.)

First we'll look at the options the program will accept. The function in Example 9-4 sets up the option parser (see "Command-line options: optparse" on page 218), but it doesn't actually parse the options.

Example 9-4. Downloading links: setting up the option parser



One of the nice things about using optparse is that the details specified for the option parser do a good job of documenting the program's behavior for the programmer, in addition to providing help strings for its user

9-5 shows the "main" Example part of the program. make command line parser to get a parser, tells it to parse the options, then checks to see that there is only one argument other than the options. If the number of nonoption arguments is not 1, the program exits with the value 2, the standard Unix value for command-line errors (as opposed to runtime errors). If there is exactly one argument, the action to be performed depends on whether the list option was specified. If it was, the program lists all the links it would download were the list not specified. Otherwise, it actually downloads them.

```
if name == ' main ':
   optparser = make command line parser()
   (options, args) = optparser.parse args()
   if len(args) != 1:
       optparser.print help()
       sys.exit(2)
                                            # command-line error
   else:
       if options.list:
           list links from url(args[0], options)
           download links from url(args[0], options)
```

Regardless of whether the program lists or downloads links, it calls get url links to extract from the file any links whose extensions appear in the list indicated by the extensions option. That function calls get url contents to read the contents of the URL, then calls extract links from string, which uses a regular expression to find all the relevant links. See Example 9-6.

Example 9-6. Downloading links: finding the links

```
def get url contents(url):
   response = urllib.request.urlopen(url)
   contents = response.read()
   response.close()
   return str(contents)
linkstring = r''([^']+?\.(exts))'|\([^\"]+?\.(exts))\""
def extract links from string(string, extensions):
   results = re.findall(linkstring.replace('exts', '|'.join(extensions)),
                         string,
                         re.M | re.S | re.I)
   return [result[1] or result[2] for result in results]
def get url links(url, options):
   return extract links from string(get url contents(url),
                                    options.extension)
```

The function get url contents implements the usual routine for reading the contents of an entire web page. Assuming link addresses are enclosed in either single or double quotes and checking the two cases separately, extract links from string replaces the substring 'exts' in the regular expression with a disjunction of the actual extensions. Only one of the two groups will match, depending on whether the address was enclosed in single or double quotes. In Chapter 8 we ignored this little problem, allowing single and double quotes to match either; here we fix that with a slightly more complex regular expression.

Listing the links just involves printing the results of get url links, and downloading the links is only a little more involved. Both are shown in Example 9-7.

The code in these examples constitutes the entire program.[†] Try it on a familiar web page to see what it does.

Web Servers

Web server technology is a mystery to many people. It conjures up images of heavy-duty hardware, special-purpose software, expert system administrators to keep things running, elaborate security precautions, and so on. Actually, though, even OS X and Windows XP can run what are called "personal web servers," and you can set them up with just a few clicks. (OS X uses the open source Apache web server that is the standard for Linux systems.)

Sockets and Servers

A web server is just one of many kind of services. You may already be aware of some others: your computer might set its time by contacting a time server; shared disks on your network are provided by file servers; and so on. In fact, time servers and file servers date back to the earliest computer networks, circa 1970. Another service you probably use is the Simple Mail Transport Protocol, which we covered in Chapter 6 (in "Sending email: smtplib" on page 224).

Such services are built on top of a low-level operating system construct called a *socket*. A socket is basically a pair of network streams managed by the operating system. A socket is identified by an address, the usual form of which is the combination of a hostname and a *port* number. The hostname can be an IP address such as 168.221.10.201, the name of a computer on the local area network (LAN), the name of an Internet host, and so on. The machine on which the program is running can be addressed with the empty string, 'localhost', or the IP address 127.0.0.1.

[†] A file containing all the code together is part of the downloadable code examples available from this book's website.

Ports with numbers below 1024 are reserved for specific services; for example, port 80 is reserved for web servers. Programs are in principle free to use port numbers from 1024 through 65535, but some ports in the range 1024 through 49151 have the status "registered"; this means that specific software products have reserved them for their own use, but unless those products are running in your program's environment no problems arise from using those port numbers.



This little discussion of sockets and servers is here to prepare you to understand what's involved in implementing programs that run on a server and respond to user requests. You are not likely to find yourself programming at this low level, but it's not inconceivable: one use for a socket-based server would be to provide to other programs on the local area network data stored across many different sources (different databases, file directories, etc.) without those programs knowing where the data is found.

Server fundamentals

A server program listens on a special server socket for connection attempts. When a client program requests a connection, the server performs the following actions:

- 1. Create a new socket with a port number generated by the operating system.
- 2. Respond to the request by telling the client the port number.
- 3. Loop reading from the socket, processing the input, and writing a response out to the socket.

The procedure is similar on the client side:

- 1. Request a connection to a server, specifying the server's hostname and the port number on which it is listening for connection attempts.
- 2. Loop writing data to the socket, reading the response, and acting on the response.

Figure 9-1 illustrates the process.

Example 9-8 shows the server code and Example 9-9 the client code for a simple directory listing server. The client accepts input from the user, in the form of a filename that can include wildcard characters; it then sends that input to the server. The server performs a glob.glob (see "Filename Expansion: fnmatch and glob" on page 232) on the pattern and returns a string consisting of all the names of the files that matched, separated by newline characters. The client reads that string and prints it.

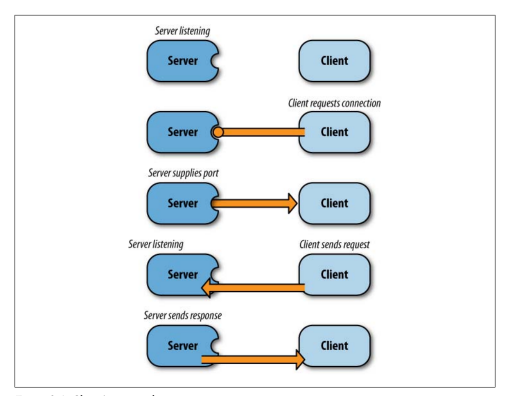


Figure 9-1. Client/server socket communication

Example 9-8. A directory listing socket server

```
import socket
import glob
HOST = socket.gethostname()
                                                          # Instead of ", allows access over LAN
PORT = 5500
                                                          # Arbitrary nonprivileged port
# listener socket
s = socket.socket(socket.AF_INET, socket.SOCK_STREAM) # standard
s.bind((HOST, PORT))
lisname = s.getsockname()
print('Listening on host', lisname[0], 'port', lisname[1])
s.listen(1)
# create actual socket
conn, addr = s.accept()
sockname = conn.getsockname()
print('Connection from', addr[0], 'on port', addr[1])
try:
    data = conn.recv(1024)
    while data:
        print(data)
```

```
conn.sendall(b'\n'.join(glob.glob(data))l
        data = conn.recv(1024)
except KeyboardInterrupt:
    pass
finally:
    try:
        conn.close()
        s.close()
    except:
        pass
Example 9-9. A directory listing socket client
import socket
HOST = socket.gethostname()
PORT = 5500
                                                              # The server's listener port number
s = socket.socket(socket.AF INET, socket.SOCK STREAM)
                                                              # standard
s.connect((HOST, PORT))
sockname = s.getsockname()
print('Connected to', sockname[0], 'on port', sockname[1])
    data = input('@ ')
    while data:
        s.send(data.encode(encoding))
        data = s.recv(1024).decode(encoding)
        print(data)
        data = input('@ ')
except (EOFError, KeyboardInterrupt):
    pass
finally:
    s.close()
```

A great many details and variations could be discussed; only the basics are demonstrated by these examples. In particular, there are important issues involved in managing multiple requests and larger amounts of transmitted data. For example, socket.sendall, used in the example's server code, keeps sending data until it has all been sent, even if it takes more than one network "packet" to transmit. The client code will have to take that into account and manage multiple packets.



As with all network communications, this example reads and writes bytes, not strings. Since the function glob.glob, used to expand wildcards on the server side, accepts bytes arguments as well as strings, the server doesn't have to convert between bytes and strings. However, socket.send does require an argument of type bytes, so the client must encode the string read from the user. The response received could be printed without being decoded (as long as the default encoding works), but it just prints as one long string; after decoding, the '\n' characters that separate the filenames are printed as newlines.

Running a server: The http.server module

Organizations often have their web servers configured so that each user has a directory from which personal pages can be served, often called *html*. Even if yours does, though, you probably don't want your programming experiments visible throughout the entire organization. Python's http.server module provides an easy way to run a simple web server for your own purposes. (Others within your LAN will also be able to access it if you want to use it for more than just experiments.)



Experimenting with running your own web server will help you better understand the fundamentals of the technology.

Example 9-10 shows the minimal setup code needed. This program will serve documents from the directory from which it was started. As it does, it will print information showing every request it gets from a client.

Example 9-10. Running a simple web server

The function is parameterized for flexibility if used with other applications, but here we just call it with the default parameters. The tuple ('', port) tells the server to listen on port. Since the official port assigned to the http protocol is 80, to access a web server that is listening on a port other than 80 you must include the port after the hostname in the URL, with a colon separating the two: http://localhost:port. This applies whether the server is running on your own computer, another computer on the LAN, or a computer somewhere on the Internet.

With the server running, enter http://localhost:8000 in the address field of a browser. You should see a listing of the files contained in the server's directory. This is the default behavior of web servers not configured otherwise for directories that don't contain a file called index.html. If one of those files is present, the browser will display that file rather than a directory listing. If a directory listing is displayed, clicking on one of the filenames will display it in the browser.



Unless the directory from which the server was started contains a file with the exact name favicon.ico, you will see repeated messages about not finding that file. A *favicon* is the tiny logo you see next to the URL in your browser's address field. Every time a browser accesses a document at a website, it also attempts to access that site's favicon.

A favicon is simply a 16×16 or 32×32 image in one of three formats: PNG, GIF, or ICO. The default name is favicon.ico, but a web page can specify a different filename and/or format (by means we won't go into here).

You might as well put one in your directory: that will stop the messages complaining about its absence and, even better, its presence in the browser will at least indicate that *something* is happening when you try to access a document even if no document appears. There's one on this book's website that you can use if you don't want to make your own. (It doesn't have to be anything special—a solid color is good enough; it just has to have one of the expected sizes and formats.)

Sometimes, especially during program development, the port the code tries to use will already be in use. This can happen because you've left a server running on that port in another window, or even because a program that aborted left the port open and the operating system didn't immediately close it. If you try the code in Example 9-10 several times in succession, interrupting it with Ctrl-Cs, you will probably see a socket.error message with the text 'Address already in use'. That's the exception raised when the program fails in its attempt to open a socket that's already open.

To deal with this problem we can extend the code so that if there's an error trying to open a port it retries with the next port number, continuing to retry until it succeeds. We'll stop before port+100, since if the code gets that far there's undoubtedly a worse problem than a conflicting port number. Example 9-11 shows the new version.

Example 9-11. Running a simple web server, finding a free port

```
def open port(hostname, startport, server class):
    for n in range(portnum, portnum+100):
        server address = ('', portnum)
            return server class(server address, handler class)
        except socket.error:
           pass
def run(portnum=8000,
        server class=http.server.HTTPServer,
        handler class=SimpleHTTPRequestHandler):
    httpd = open port('', 8000, server class)
    hostname, portnumber = httpd.socket.getsockname()
    print('Serving HTTP documents on {}, port {}...
         .format(hostname, portnumber))
    httpd.serve forever()
```

CGI

From the earliest days of web servers, requests submitted through web pages have used a mechanism called the *Common Gateway Interface* (CGI). There are more sophisticated mechanisms now, but CGI is still widely used. The CGI mechanism is nothing more than a standardized way of passing a URL request and its arguments to a program. After receiving the request, the program does its work and outputs HTML that gets sent back to the browser for display. The Python module that supports this mechanism is cgi.

As we saw at the beginning of this chapter, a URL often represents a request, rather than a specific document. The address part of a request identifies a program to be executed. A question mark identifies the beginning of the part of the URL containing the arguments of the request and their values. These arguments are the equivalent of keyword arguments in Python function calls, though they are not in any way specific to Python.

Serving CGI requests

A web server must be configured to execute CGI programs, or else a URL for a CGI program will simply display its text in the browser. Most web servers are configured to run CGI scripts. There are two ways they can distinguish CGI scripts from regular web pages: by their extension or by their location within the file hierarchy from which the server serves pages. Organizations that configure their web servers so that each user has a designated directory from which web pages may be served generally also configure things so that a subdirectory of that directory, generally called either *cgi-bin* or *cgi*, can contain CGI scripts.

If your organization has its web server configured appropriately, or you have a personal web server running on your computer that you can configure, you can use that server for experimenting with CGI scripts. In general, though, it is easier to do your experiments and development using Python's own simple server, as was shown in Example 9-10. The code in that example, however, only serves documents. To enable it to serve CGI requests too, the class CGIHTTPRequestHandler must be used in place of SimpleHTTPRequestServer, as shown in Example 9-12.

The default directories that contain programs CGIHTTPRequestHandler can run are listed in CGIHTTPRequestHandler.cgi_directories. The default value is ['/cgi-bin', '/htbin'], but your program could change that (perhaps adding '/cgi' to the list, as is done in Example 9-12). These subdirectories are relative to the directory from which you start the server—if you start the server from <code>/home/me/python/web</code>, that's what the server will interpret / to mean.

```
Example 9-12. Running a CGI server
```

```
http.server.CGIHTTPRequestHandler.cgi_directories.append('/cgi')
def open port(hostname, startport, server class):
```

```
for n in range(portnum, portnum+80):
        server address = ('', portnum)
        try:
            return server class(server address, handler class)
        except socket.error:
           pass
def run(portnum=8000.
        server class=http.server.HTTPServer,
        handler_class=http.server.CGIHTTPRequestHandler):
   httpd = open_port('', 8000, server_class)
   hostname, portnumber = httpd.socket.getsockname()
   print("Serving HTTP, including CGIs, on",
         hostname,
          "port",
         portnumber, "...")
   print('Serving HTTP documents and CGI requests on {}, port {}...'
         .format(hostname, portnumbe))
    print("CGI directories are:",
         http.server.SimpleHTTPRequestHandler.cgi directories)
   httpd.serve forever()
```

Documents can go in the same directory as the server, but executables must be in one of the subdirectories listed in the value of CGIHTTPRequestHandler.cgi directories. Note that CGIHTTPRequestHandler can distinguish CGI scripts from documents only by their being located in one of these directories; it does not support extension-based determination the way full-powered web servers do.

Setting up CGI

When you first start working with CGI scripts, there are many mysterious things that can go wrong. Some of them result in completely blank web pages, with no hint of the nature of the problem. Sometimes what you get is a listing of the program instead of the result of its execution. Some of the problems are caused by various aspects of your CGI setup. Before trying to use your own code, you should take advantage of the cgi module's test function, which will send the browser a large amount of information about the CGI environment. Example 9-13 shows a complete CGI program using cgi.test. If cgi.test doesn't work, you haven't set things up entirely correctly.

```
Example 9-13. Running cgi.test
import cgi
cgi.test()
```

Run the program from Example 9-12 in a command window. It will run continuously until you interrupt it. Note the port it says it is listening on when it starts. Put the program shown in Example 9-13 in one of the subdirectories the program shows, naming it *cgitest.py*. On an operating system that supports "execute" permissions, change (with the chmod command-line command) the permissions of the file so that everyone can read and execute it (e.g., 755 on Unix-based systems).

Now, enter the following URL into your browser's address field, replacing *port* with the number of the port on which the server program is listening and *cgi* with the name of the directory in which you put *cgitest.py*:

```
http://localhost:port/cgi/cgitest.py
```

Your browser should display a long list of details of the environment in which the program ran.

CGI script arguments and responses

You won't learn much from cgi.test, except whether your CGI server environment is working and that you have your CGI script in the right directory with the right permissions. You will not see two lines that cgi.test prints first that are necessary to inform the browser that the program is sending it HTML to display. All CGI scripts must print these two lines before any other output:

Content-Type: text/html

<!-- the previous line must be blank; this line is just an HTML comment -->

Like many of the Python library's more frequently used modules, cgi provides several levels of interfaces. The simplest will suffice for most routine CGI scripting. The CGI Script template presented here shows the general outline of a CGI program. The program starts with the usual "shebang" line that specifies the program used to run the script. (This takes the place of typing python3 and the name of the program at the command line.) To be run as a CGI script, the file must also be readable and executable by everyone, so set the file's permissions accordingly.

Because the shebang line specifies that this is a Python file, it does not need the .py extension to run. There's no need for users to be distracted by extensions like that, so it's good form to remove the extension from the filename. However, the extension is still needed so that operating systems, editors, and IDEs recognize it is a Python file when you open it for editing. The way around this conflict is to keep the .py extension on the file you are developing and create a link ("alias" or "shortcut") to it that does not have the extension. This also allows you to keep your files in a different directory than the one containing your CGI scripts—the link can be made from the file to your CGI directory.

When a CGI script is executed through a web server, everything it prints to sys.stdout is sent to the user's browser for display. (The newlines printed in the template are just for readability, for viewing a page's HTML "source" in a browser's "view source" window or the output when executing the program from the command line; in most contexts browsers ignore whitespace between the end of one HTML tag and the beginning of the next.)

TEMPLATE

Structure of a CGI Script

```
import cgi
import cgitb
cgitb.enable()
def respond():
   print(
                             # the second line printed must be blank
'''Content-Type: text/html
<html>\n<head>\n<title>title</title>\n</head>\n<body>
   args = cgi.FieldStorage()
    ... access args using getfirst and getlist ...
    print the HTML body
    print('</body>\n</html>')
if __name__ == '__main__':
    respond()
```

At the beginning the template initializes a very useful tool for debugging CGI scripts, which will normally be removed for a production script: importing and enabling cgitb arranges for nicely formatted runtime error reports to be displayed in the browser. It does not catch syntax errors, though—if your program has syntax errors, all you will see in the browser is a blank page. (You will see syntax errors reported in the window in which the server is running.)

The crux of Python's CGI module is the FieldStorage class. When an instance of the class is created, it parses the URL's query string and stores its arguments and values. Subsequently, the value of a query argument can be obtained by accessing the FieldStorage object as if it were a dictionary.

Because a request may have multiple values for the same argument, FieldStorage must support both single- and multiple-value arguments. (Even if your program expects just one value for an argument, a user might type in a URL that has more than one.) Dealing with this can get awkward, so FieldStorage provides two convenience methods: getfirst(arqname) to return one value, and getlist(arqname) to get a list of however many were present in the request. You can provide a second argument to getfirst (default None) that specifies the value to return if the request URL had no value for the argument. Normally it's better to obtain argument values this way rather than by accessing the values directly with dictionary notation.

The mechanism by which a web server passes arguments to a CGI script is really quite simple: all it does is set the environment variable QUERY STRING to the part of the URL that follows the question mark. (There is another, more elaborate way of passing arguments that doesn't put them in the query string; getting the arguments in that case is a bit more complicated.) When a FieldStorage object is created, it checks whether the os.environ dictionary contains the key 'QUERY_STRING'. If the key is present, its value is used to populate the FieldStorage object. If the environment does not contain 'QUERY_STRING', the FieldStorage object next looks for a command-line argument. If sys.argv[1] is present, it uses that as the query string.



This means that you can test your CGI scripts by giving them a query string as a command-line argument. You don't need the whole URL, just the part after the question mark. This is especially useful for eliminating syntax errors before trying to run the script from a browser.

Let's fill in the CGI Script template with enough details to "echo" the arguments and values supplied in a URL. Example 9-14 shows what this would look like. In general, CGI scripts ignore arguments with unrecognized names; to demonstrate this usage, the program will only look for one argument named enzyme and any number of arguments named sequence.

Example 9-14. A CGI echo script

```
import cgi
import cgitb
cgitb.enable()

def respond():
    print("Content-Type: text/html")
    print()
    print("<html>\n<head>\n<title>title>\title>\n</head>\n<body>")
    args = cgi.FieldStorage()

# completing the template:
    print('Enzyme:', args.getfirst('enzyme'), '')
    print('Sequences:', args.getlist('sequence'), '')
    print("</body>\n</html>")

if __name__ == '__main__':
    respond()
```

If you have everything set up correctly, typing the following URL (together on one line, not with the line break):

```
http://localhost:8000/cgi/cgi_echo.py?enzyme='EcoRI'&
sequence=CCCC&sequence=CGG&sequence=GGGG
```

into a browser should result in a web page that displays the following, unformatted:

```
Enzyme: 'EcoRI'
Sequences: ['CCCC', 'CCGG', 'GGGG']
```

Simple Web Applications

We'll close this section on web servers with a few real examples. The first is a straightforward application of the CGI techniques discussed earlier. The second will introduce some new information about creating web pages with simple forms and using those forms to submit requests to CGI scripts.

Using CGI scripts

This section's example is a CGI program to search Rebase data for all enzymes that recognize a particular site, specified as a site query argument in a URL. We'll assume that we have a datafile called *bionet.table* (bionet being one of the Rebase file formats), as written by the write table function of Example 4-41 (see "Step 5" on page 159). Each line of that file contains the name of an enzyme, a tab, and a recognition sequence. Some of those recognition sequences contain a caret (^) indicating where the enzyme cuts the sequence, but we'll ignore those for this program.

Reading this file into a dictionary is simple, but there's a twist: we need to reverse the table. Instead of looking up recognition sites by enzyme, we want to find all the enzymes that recognize a particular site. Example 9-15 shows how to do this.

Example 9-15. Reading Rebase data into a reversed dictionary

```
def read table(filename):
   table = {}
   with open(filename) as fil:
        for line in fil:
            enzyme, sequence = line.split()
            sequence = sequence.replace('^', '')
                                                     # ignore cut sites
            table.get(sequence, []).add(enzyme)
   return table
```

To generate the HTML for the CGI response, we are going to take advantage of the template facility in the string module (see "String Utilities: string" on page 240). This makes it easy to specify a large chunk of text that has only a few variable parts, which is just what we need (Example 9-16). In fact, we'll use a series of templates.

Example 9-16. The main template for the recognition site script

```
html template = string.Template(
                                                        # second line is requisite empty line
    '''Content-Type: text/html
<head>
<title>Restriction Enzyme Search</title>
</head>
<body>
<h2>Restriction Enzyme Search</h2>
$response
</body>
</html>
''')
```

There are three conditions the program must handle:

- 1. The URL did not include a site argument.
- 2. There are no enzymes that recognize the site.
- 3. There are one or more enzymes that recognize the site.

The program's responses for the three cases will be:

- 1. Display an error message.
- 2. Display a message that no matching enzymes were found.
- 3. Display an ordered (numbered) list showing the enzymes that recognize the specified site.

For each case the program will substitute appropriate HTML for the \$reponse placeholder of html template from Example 9-16. For the second and third cases the program uses templates to generate the response, as shown in Example 9-17; for the first case it just uses a plain string.

Example 9-17. Response Templates for the recognition site script

```
none recognized template = string.Template(
    '''<i>No enzymes recognize <b>$seq</b>.</i>\n''')
response template = string.Template(
    '''Enzyme(s) recognizing <b>$seq</b> are:
<01>
$items
''')
```

Example 9-18 shows the code that gets the script's one argument and produces the HTML to return to the browser. It initializes a dictionary for use with the main template, then for each of the three cases it sets the value of its 'response' entry appropriately before performing a substitution.

Example 9-18. Code for the recognition site script

```
def make html body(table, seq):
   subst = {'seq': seq, 'items': [], 'response': ''}
   if not seq:
                                                        # case 1 -- no value for sea
        subst['response'] = 'No value for site in query arguments'
        seq = seq.upper()
                                                        # doesn't change value in subst dict
        if seq not in table:
                                                        # case 2 -- no match found
            subst['response'] = \
                none recognized template.substitute({'seq': seq})
        else:
                                                        # case 3 -- found, make list items
            items = '\n'.join(['' + enzyme + ''
                                for enzyme in table[seq]])
            subst['response'] = \
                response template.substitute({'seq': seq, 'items': items})
   return subst
```

```
def print response(table, seq):
   print(html template.substitute(make html body(table, seq)))
def respond():
   print response(read table('data/Rebase/bionet.table'),
                  cgi.FieldStorage().getfirst('site'))
if name == " main ":
   respond()
```

As described earlier, you can test this script from the command line. For example:

```
% python3 enzymes for site.py site=cccggg
```

The output from this example should be HTML that lists three enzymes recognizing 'CCCGGG'. (The script uses string.upper to make the sequence lookup case-insensitive.) If you get a runtime error complaining that the file bionet.table cannot be found, make sure that that file—or a link to it—is present in the directory from which the program runs, whether from the command line or through CGI.



If you run the CGI program from a browser through a full-scale web server rather than using http.CGIHTTPRequestHandler, the datafile's permissions must be set to allow it to be "readable" by everyone, since scripts will not be run under your username.

HTML forms with CGI scripts

Although a URL for a request can be constructed programmatically, as in Example 9-1, requests are usually generated from web pages. HTML forms are the original method for doing this, and while old and simplistic, they are still a reasonable approach for simple web pages. More recent advances in client-side technologies are far beyond the scope of this book. The purpose of introducing HTML forms here is twofold:

- 1. It's easy to put together a simple interface with them, since the user's browser does most of the work.
- 2. It demonstrates the round-trip interactions involved in *client/server* technologies, of which web forms together with CGI scripts are an important example.

An HTML form is specified by a form tag. Any HTML tags and text can appear between the opening <form> and closing </form> tags. Certain tags designate form controls, which have fairly standardized appearance and behavior.

Generally, each control tag has a name attribute, which is used as the name of the parameter in any HTTP request the form submits. The form tag itself has an action attribute that specifies the URL of the program that will run when the form is submitted. (Frequently this is a relative URL—i.e., relative to the location of the page containing the form—though that is not a requirement.) When the request is submitted you will see a new URL in your browser's address bar that includes the path to the CGI script, a question mark, and *name=value* pairs for each field in the form.



Being able to see the request generated by submitting the form is a very helpful aid to debugging and learning more about CGI.

Client-side scripts (usually in JavaScript) may be attached to various events related to controls. Client-side scripts can also change the state of form controls. We will ignore client-side programming here and talk only about the default behavior of each kind of control and the interactions users have with them.

When a form is submitted, the value of a control is paired with the tag's name attribute to form the *name=value* pairs in the URL. The value is obtained from the state of the control in the browser—control tags do not need value attributes. However, if the form's HTML provides a value attribute for a control tag, that value will be used to initialize the state of that control. The kinds of tags that constitute forms are:

button

There are three types of button controls, specified by the value of the tag's type attribute:

submit

Clicking a submit button causes the form to be submitted—i.e., the names and values of the form's controls are combined with the form's action URL to create a request, which is then submitted to the web server (normally the server from which the form was obtained in the first place).

reset

Clicking a reset button restores all the controls to their initial state.

push

Push buttons have no default behavior; they are used together with scripts, so we won't be discussing them any further here.

input

Input controls include the following, specified by the value of the tag's type attribute:

checkbox

On/off switches that the user can toggle. Several checkboxes in a form may share the same name attribute value, allowing multiple values to be selected.

radio

Like checkboxes, but they have a different appearance and normally share a name attribute value. The browser ensures that only one is selected at any time.

text

A single-line area where the user can type text that becomes the value of the control.

file

Opens a file selection dialog for uploading files.

textarea

A multi-line input control (text box).

select

A drop-down menu, each value of which is specified by an option tag.

Example 9-19 shows the HTML for a very simple web page. It has a form containing three elements: a label, an input box, and a submit button. The intent of the form is to generate a request to search the Rebase data loaded by the program shown in Examples 4-37 through 4-41.

Example 9-19. A simple HTML form for a restriction site query

```
<!-- begin HTML -->
  <head>
                                                              <!-- begin HEAD -->
    <title>Restriction Enzyme Search </title>
                                                              <!-- window title -->
  </head>
                                                              <!-- end HEAD -->
                                                              <!-- begin BODY -->
  <body>
    <h2>Restriction Enzyme Search</h2>
                                                              <!-- heading 2 -->
    <form action='cgi/enzymes for site'>
                                                              <!-- begin FORM -->
    <!-- action attribute specifies CGI script to run on submit -->
      <label>Recognition Site:</label>
                                                              <!-- just text -->
      <input type='text' name='site' size='25'>
                                                              <!-- 25 char. input -->
      <button type='submit'>Find</button>
                                                              <!-- submit button -->
    </form>
                                                              <!-- end of FORM -->
  </body>
                                                              <!-- end of BODY -->
</html>
                                                              <!-- end of HTML -->
```

Figure 9-2 shows how a browser would display the form shown in Example 9-19. (Of course, the actual fonts and possibly other details will differ according to which browser is used and how its preferences are set.)

Restriction Enzyme Search			
Recognition Site:	Find		

Figure 9-2. The restriction site query form

Suppose the URL for the HTML page displayed in Figure 9-2 is:

```
http://localhost:8000/enzymes for site.html
```

If the user enters the sequence CCGG in the text input field and presses the submit button, the browser will submit the following request:

```
http://localhost:8000/cgi/enzymes for site?site=CCGG
```

The form's action attribute does not begin with an http:, so its path uses the same host as the HTML file. It doesn't begin with a slash, so the path is relative to the directory from which the server is running. Together that gives the partial URL http://local host:8000/. The rest of the path, cgi/enzymes_for_size, is the value of the submit attribute of the <form> tag. When a form like this is submitted, the name and value of each of its input components is added to the URL, following a question mark. The name and value are separated by an equals sign, and if there are more than one the pairs are separated by ampersands.

The result will be something like what is shown in Figure 9-3.

Restriction Enzyme Search

Enzyme(s) recognizing AATT are:

- 1. TasI
- 2. MluCI
- 3. Sse9I
- 4. TspEI
- 5. TseCI
- 6. Tsp509I

Figure 9-3. The restriction site query results

There's one more thing to take care of. The HTML produced by the preceding code does not include a form. Once the user has submitted one query, the web page returned can't be used to submit another one. True, the user could use the browser's back button, but for various reasons that is not a particularly good solution. It would be better to return the original form *plus* the results. All we'd need to do to implement this is extend the main HTML template to incorporate the HTML from the original form into the HTML for the response. Example 9-20 shows the new version of the HTML template originally defined in Example 9-16.

```
Example 9-20. Returning a form with CGI results
```

```
html_template = string.Template(
    '''Content-Type: text/html

<head>
    <title>Restriction Enzyme Search</title>
</head>
<body>

    <form action='enzymes for_site'>
          <label>Recognition Site:</label>
          <input type='text' name='site' size='25' value=$seq>
          <button type='submit'>Find</button>
```

```
</form>
$response
  </body>
</html>
''')
```

Now, the user will see the results below the input area and will be able to use the form again. In addition, the previous entry in the input form will still be there, ready for editing in case the user wants to change it slightly. Figure 9-4 shows the results of the same query as Figure 9-3 after changing the HTML template as shown in Example 9-20.

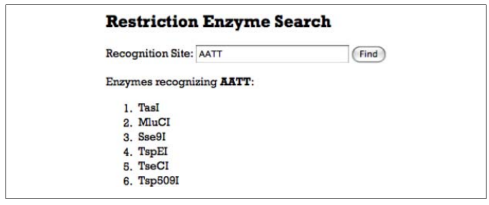


Figure 9-4. The restriction site query results with a form

We no longer even need the original HTML file now. We can just enter the query URL with no arguments to get the empty form:

http://localhost:8000/cgi/enzymes for site



The cgi/component is not part of the action attribute of the generated HTML because the path will be relative to the directory where the CGI script that generated it is found.

Tips, Traps, and Tracebacks

Entrez Programming Utilities

The Entrez Programming Utilities (E-Utilities) provide uniform access to many of the Entrez databases. They are accessed through HTTP queries, but their parameters and responses are designed for programmatic use, as opposed to HTML forms and web pages marked up with formatting details. One of the query parameters specifies the desired form of output: text, XML, etc. The starting point for their documentation is http://eutils.ncbi.nlm.nih.gov/entrez/query/static/eutils_help.html. That page provides links to the documentation of each of the tools and a short course describing how to use them to construct data pipelines (http://www.ncbi.nlm.nih.gov/bookshelf/br.fcgi ?book=coursework&part=eutils). Perhaps the most useful query in the context of the kinds of data access shown in this book is Efetch, specifically in its use with "Sequence and other Molecular Biology Databases," as described at http://eutils.ncbi.nlm.nih.gov/ corehtml/query/static/efetchseq help.html. A complete list of the software tools available from the NCBI website is at http://www.ncbi.nlm.nih.gov/guide/data-software.

Tips

Web pages

• Put the following function, which we saw earlier, in a file of Python utilities you maintain, and import it into the files you are working on. (For a module like this it does make sense to use the from ... import * form of the import statement.) You can call this function from within Python to open the documentation of a given module. You'll probably want to give it a shorter name, but there's no need to change the name of the function itself. Instead, simply assign the name you want to browse module doc:

```
def browse module doc(module name):
     webbrowser.open('http://docs.python.org/3.1/library/' +
                      module name + '.html')
mdoc = browse module doc
>>> mdoc('re')
                         # browse re module doc
>>> mdoc('os.path')
                         # browse os.path module doc
>>> mdoc('modules')
                         # browse modules index
>>> mdoc('index')
                         # browse library index
```

- Like Python strings, HTML attribute values may use single or double quotes. Mixtures of single and double quotes can cause a great deal of confusion and frustration when you are constructing HTML strings, matching HTML text with regular expressions, or using ordinary string facilities to take apart HTML. If your program is constructing HTML, you can choose which quotes to use where, and you should make a point of using either single quotes for Python strings and double quotes for HTML tag attributes, or vice versa. If your program is analyzing HTML obtained from elsewhere, and that HTML uses either single or double quotes consistently, use the other kind for your strings. Of course, if you always use triple quotes in these situations—even for one-line strings—you will avoid such problems entirely.
- It is very tricky to develop and debug both web interactions and program actions together. Whenever you do any kind of web programming, develop the code first using data from a locally stored file. (In particular, if the data is supposed to be obtained by a query, perform the web query manually and save the results to a text file.) Once your code is working, add the actual web interaction.

Debugging CGI scripts

- In general, you can run your CGI script from the command line, giving it a single argument that contains arguments the way they would appear in a query following its question mark: in the form of *name=value* pairs separated by ampersands. This can be very helpful for debugging. (Usually you'll have to quote the argument on the command line, especially when there is more than one value, since most command-line shells treat ampersands specially.)
- Importing your script into the interpreter and running pieces of it manually is another effective debugging technique.
- Always import and include cgitb.enable in your code so that you get its elaborate runtime error messages while developing CGI scripts.
- Syntax errors in CGI will show up in the window from which you are running one of the http.server module's servers.

HTTP servers

- The http.server module's servers print every request they get to the window in which they are running.
- You can stop the favicon messages by installing a favicon.ico file (see "Running a server: The http.server module" on page 341) in the directories from which you run web servers (Python's or otherwise).
- Use links (aliases) to .py CGI files, omitting the file extension. This is a form of information hiding. (This may not work in Windows environments, where the extension is the only way to tell the system what program to use to execute a file.) Keep the following points in mind:
 - —You want the URL submitted to look like a query, not a filename. The implementation of the query as a Python script is a detail that should not be exposed to the outside world. (Don't you get a bit irritated or confused when you see .php or .pl or even .cgi extensions in URLs?)
 - —Because the alias separates the query name from the actual file that implements it, you can change the path to the actual file and re-create the alias or link to the new path without having to change anything in the HTML files that use the query in form actions.
- Give your queries (the aliases or links to your .py files, or the .py files themselves if you aren't using aliases or links) descriptive but not enormously long names. Users may read the query URLs in their browsers' address bars, bookmark them, etc. Later, they may search their history or bookmarks for a URL and should be able to find it by recalling a natural syllable in the query name.
- Likewise, keep the paths of your query aliases relatively short, even if the actual files are buried deep in your file hierarchy.

• When you're running an http.SimpleHTTPServer or http.CGIHTTPServer from a command-line Python program, syntax errors will appear in the window from which it was started. Tracebacks will not. In order to view tracebacks you must include the following lines in your program:

```
import cgitb
cgitb.enable()
```

Those tracebacks will appear (in dramatic color) in the web browser. Obviously, you would not want to enable this when people are actually using your CGI script.

Traps

- All network programming protocols return bytes objects. You must call the decode method on bytes objects you want converted to strings. You can specify the encoding to use as an argument; otherwise, the default decoding (usually the local variant of ASCII) will be used.
- Web servers execute CGI scripts as separate executables with no shell environment. Therefore, if using a "shebang" line it must specify the absolute path to the Python executable, rather than using /usr/bin/env.
- For a file to be executed as a CGI script using an http.server module server:
 - —You must use http.CGIHTTPServer: it both serves documents and executes CGI scripts, whereas http.CGIHTTPServer only serves documents.
 - —The script file must be executable.
 - The http.CGIHTTPServer recognizes a request as a CGI script simply by checking the path from the directory in which the server was started to the directory of the script. Extensions (or lack thereof) have no effect. The subdirectory can be several levels down.
- Absolute and relative URLs in an HTML file are treated very differently. Absolute URLs begin with a slash: their paths begin at the directory from which the web server is running. Relative URLs do not have a leading slash; their paths begin at the directory from which the web page was obtained.
- Running scripts through a full-featured web server rather than using one from Python's http.server module introduces further constraints. The exact details depend on the site's configuration, but the following are likely to be issues. Many are the result of the server's executing scripts under a special username reserved for the server. The considerations in this list were much more of an issue in earlier days of organization web servers, but you should be aware of them, especially if your organization may be using default configurations that retain some of these restrictions and requirements:
 - —Scripts must be readable and executable by "anyone."
 - —Datafiles accessed by scripts must be readable by "anyone."

- —Directories on the path from a script's directory to any datafile it uses must be navigable (probably readable and executable) by "anyone."
- —Whether a file is interpreted as a script to execute may depend on the name of the directory in which it is found, its extension, or a combination of the two. (If only certain extensions are allowed, you may have to use a .cgi extension in the alias or links to your actual .py files.)
- —It may be necessary to place scripts and HTML files in designated directories under your home directory—html, for example.
- —Your scripts may not be accessible from the Internet, depending on the site's configuration, firewall settings, etc.
- —Links or aliases from within your designated HTML and CGI subdirectories to files in directories outside of those directory trees may not be allowed, or they may work for HTML but not for CGI.

Tracebacks

Here are some error messages you are likely to encounter while doing web programming with Python's libraries:

socket.error: Address already in use

An attempt has been made to start a server on a port that is already being used; typically, this will happen when a server you started has not stopped and released its port—either kill the process manually or start a new server using a different port number.

TypeError: can't use a string pattern on a bytes-like object

You matched a regular expression string of type str against bytes returned from urllib.request.urlopen; convert the bytes to a string by calling str, with an encoding argument if necessary, or call the decode method on the bytes, giving it an *encoding* argument if necessary.

urllib.error.HTTPError 400: Bad Request

An error has been returned from a web server in response to a call to urllib.request.open or the like.

urllib.error.URLError: <urlopen error [Errno 8] nodename nor servname provided, or not known>

In a call to urllib.request.open, the URL didn't contain a hostname, or your computer is not connected to the Internet.

Relational Databases

Most generally, a *database* is simply an organized collection of data. The data can be stored in memory, in files in one of your directories, in publicly accessible files, or in files on a computer acting as a *database server*. Like a web server, a database server stands between client programs and the actual data, so that applications never directly access the files storing the data.

There isn't necessarily anything special about database files, other than the structure given to their data by the applications that access them. Spreadsheets have long been used informally for recording and manipulating data in files. Desktop office application suites generally include a database component that offers capabilities beyond those of spreadsheets. These database components allow you to access data selectively instead of having to read and write entire spreadsheet documents. They also provide mechanisms like pattern-based searching that can be used to construct desktop applications.

Database servers, in turn, are far more powerful and can hold far more data than either spreadsheets or desktop office suite database components. You may have heard of commercial database systems such as Sybase and Oracle and open source systems such as PostgreSQL and MySQL. Database servers use highly specialized file formats that support capabilities such as concurrent access by multiple users, restrictions on access to data, a variety of efficiency techniques, automatic backup, and a wide range of other features necessary for group- and enterprise-scale data management. Client applications access data on these servers through a standardized language known as the *Structured Query Language* (SQL).

Most database server technology is based on a *relational data model*. The theory of relational data models is a rather complex subject. Fortunately, we can use the less formal terminology of SQL, the standardized language used with relational databases.* This section presents a very basic introduction to the relational model and SQL. Next, we'll explore an example of designing and loading data into a database.

^{*} As an acronym for Structured Query Language, "SQL" should be pronounced "ess cue ell," though many people pronounce it like the word "sequel."

Once we have data in a database, we can look at some examples of database queries. The chapter culminates with the development of a web page and a CGI script that allow the database to be queried through a form in a browser.

Representation in Relational Databases

A full introduction to relational database design would more comfortably occupy an entire book than a section of a chapter. The goal here is to give you enough information to get you started using relational databases. We will cover the essentials and demonstrate them with an example.

Database Tables

A relational database stores data in tables that, like spreadsheets, are organized into rows and columns. Each row represents an individual entity (a neutral term used to avoid committing to a particular interpretation of what the data in the row means). Each column represents an attribute of the entities in the tables. In Python terms, a table is like a class (type), a row is like an object, and a column is like an attribute, with each column of a row holding the value of the corresponding attribute.

As in Python dictionaries, the entries in relational database tables are identified by keys. A key for a table's entries may comprise the value of a single column or the values of several columns. Because a table's key identifies an individual row, each row's key must be distinct from the keys of every other row in that table.

A table may have more than one key, since there may be more than one column or combination of columns that reliably distinguishes each row from every other. Usually, however, one of the keys is designated as the *primary key*. This is not a requirement of the relational data model, but a pragmatic aspect of the way database systems define and manage tables. Database systems also offer a mechanism by which an integervalued column can be designated as a primary key with values automatically generated in sequence.

Avoiding duplication of values

One important principle of good database design is that combinations of column values should not appear in more than one row of a table. To give a minimal example, suppose we write down the four ways to refer to an amino acid, as shown in Table 10-1. This violates the rule, because each combination of values for Code, Short, and Name occurs in more than one row.

Table 10-1. A minimal example of column dependency

Codon	Code	Short	Name
UUU	F	Phe	Phenylalanine
UUC	F	Phe	Phenylalanine
UUA	L	Leu	Leucine
UUG	L	Leu	Leucine
UCU	S	Ser	Serine
UCC	S	Ser	Serine
UCA	S	Ser	Serine
UCG	S	Ser	Serine

Bad things happen when this rule is violated. For example, "phenylalanine" may be mistakenly spelled differently in the two rows in which it appears. That would mean that "UUU" and "UUC" would refer to different amino acid names even though they refer to the same amino acid code and short names, creating a serious inconsistency in the data. Another problem is that repeating combinations of column values causes the table to take up more space than is necessary.

The way to fix these kinds of problems is to create a second table to contain just a single copy of the repetitive information, as shown in Table 10-2. Any of the three columns in this table can be used as a key, because given the value of one of the columns, the values of the other two are uniquely defined.

Table 10-2. A database table for amino acid codes and names

Code	Short	Name
F	Phe	Phenylalanine
L	Leu	Leucine
S	Ser	Serine

Table 10-1 can now be revised by dropping two of the redundant columns, as shown in Table 10-3.

Table 10-3. A revised codon-amino acid table

Codon	Code	
UUU	F	
UUC	F	
UUA	L	
UUG	L	

Codon	Code
UCU	S
UCC	S
UCA	S
UCG	S

Given these two tables, here are the steps for obtaining the full name of an amino acid given a codon:

- 1. The row with the specified codon as the value of its Codon column is located in the codons table.
- 2. The value of the Code column of that row is used as a key to the amino acids table.
- 3. Finding the row with that value in its Code column, the value of the Name column is obtained.

In the vocabulary of dictionaries, the value of an entry in the first dictionary is used as the key to access a value in the second dictionary. The technical term for one or more columns of one table that are used to look up a row in another table is a *foreign key* foreign because it's a key in a different table.

Representation of relationships

Together, Tables 10-2 and 10-3 define a relationship between codons and amino acids. This is called a *one-many* or 1-N relationship, because for every one entry in the amino acid table there are potentially many (N) entries in the codon table. Occasionally a database will have a one-one or 1-1 relationship, where each entry in one table corresponds to at most one entry in the other. More common than 1-1, but less common than 1-N, are relationships in which for each entry in one table there can be many in the other, and vice versa. This kind of relationship is called a many-many or M-N relationship.



Strange as it may sound, the term "relationship" is not what the word "relational" refers to in the terms "relational database" or "relational data model." "Relation" is a mathematical term akin to "function," in the sense of mapping a domain into a range. "Relationship" is a design and implementation term signifying an interconnection among objects.

M-N relationships require special handling. Consider a table representing researchers (Table 10-4) and another representing institutions (Table 10-5). The primary key for the Researcher table will be the researcher's Social Security number or equivalent ID. In the Institution table we'll make the simplifying assumption that no two institutions have the same name, so we can use the name as the primary key. (If two institutions

did have the same name, we could use a combination of the institution's name and state as the primary key, or we could use something more definitive, such as the institution's government tax ID.)

Table 10-4. A table of information about researchers

ID	First	MI	Last	Position	Institution
111-22-3333	Blue	J	Bird	Professor	Some Univ.
444-55-6666	Kay	С	Jones	Student	Another Univ.
777-88-9999	Elm		Oh	Postdoc	Some Univ.

Table 10-5. A table of information about institutions

Name	Street	City	State	Country
Some Univ.	1 A St.	Oneville	MA	USA
Another Univ.	20 B St.	Twotown	NY	USA
Corporate R&D	9 Ave. Q	Oneville	MA	USA

So far we have the same kind of 1-N relationship we had in the previous example: N researchers for any institution, one institution for any researcher. Now let's make Blue J. Bird a consultant at Corporate R&D, as shown in Table 10-6.

Table 10-6. Adding another entry for a researcher

ID	First	MI	Last	Position	Institution
111-22-3333	Blue	J	Bird	Professor	Some Univ.
444-55-6666	Kay	С	Jones	Student	Another Univ.
777-88-9999	Elm		0h	Postdoc	Some Univ.
111-22-3333	Blue	J	Bird	Consultant	Corporate R&D

Now we have a problem: rows 1 and 4 have the same primary key—the ID—which violates the requirement that no two rows have the same value for a key. The problem here is that an M-N relationship is essentially two 1-N relationships: one researcher can have multiple positions and, of course, one institution can have multiple researchers. However, in a 1-N relationship only one of the tables can have rows that repeat values for the foreign key to the other table.

M-N relationships must be represented in a separate table whose rows contain foreign keys to each of the other two tables. First we'll create a table called "Appointment." We'll add to it a column for the foreign key from each of the two tables involved in the relationship. The primary key for this table will be the combination of the values in those columns. Table 10-7 illustrates this initial step. Note that no two rows have the same value for the combination of researcher and institution.

Table 10-7. Creating an M-N Appointment table

Researcher	Institution
111-22-3333	Some Univ.
444-55-6666	Another Univ.
777-88-9999	Some Univ.
111-22-3333	Corporate R&D

Next, we'll remove the foreign key column from the Researcher table, as well as the columns that differ between the rows with the same ID. We'll then add those columns to the new table. In this example the only column other than the foreign key is the Position column. Moving the Position column to the Appointment table makes sense, because a position is associated with the combination of researcher and institution: a person isn't a professor, but rather a professor at a particular institution. Table 10-8 shows the modified Appointment table and Table 10-9 the modified Researcher table.

Table 10-8. Adding a column to the Appointment table

Researcher	Institution	Position
111-22-3333	Some Univ.	Professor
444-55-6666	Another Univ.	Student
777-88-9999	Some Univ.	Postdoc
111-22-3333	Corporate R&D	Consultant

Table 10-9. Adding a column to the Researcher table

Researcher	First	MI	Last
111-22-3333	Blue	J	Bird
444-55-6666	Kay	C	Jones
777-88-9999	Elm		Oh

Database definition

The table definitions of a relational database constitute its schema. Many formal notations and diagramming techniques exist for specifying complex schemas, but for simple databases it is often sufficient just to list the table details in text format. Specifically:

- The names of the database's tables
- The name of each column of each table
- The type of value contained in each column of each table
- For each column of each table, whether its values can be null
- The column(s) of each table forming the key by which its entries can be referenced

- The primary key of each table
- The columns of each table having values that are keys in another table (the foreign keys)

A database containing both the amino acid tables and the researcher tables could be described textually as shown in Table 10-10. (The asterisks denote primary keys; the type char represents a fixed number of characters, while varchar represents a variable number of characters up to the specified limit.)

Table 10-10. An informal database schema

Table	Column	Туре	Null?	Foreign key
AminoAcid	*Code	char(1)	N	
	Short	char(3)	N	
	Name	char(13)	N	
Codon	*Codon	char(3)	N	
	Code	char(1)	N	AminoAcid.Code
Researcher	*ID	integer	N	
	First	varchar(255)	N	
	MI	char(3)		
	Last	varchar(255)	N	
Institution	*Name	varchar(255)	N	
	Street	varchar(255)		
	City	varchar(255)	N	
	State	char(3)	N	
	Country	char(5)		
Appointment	*ID	text	N	Researcher.ID
	*Institution	text	N	Institution.Name
	Position	text		

A Restriction Enzyme Database

In this section we're going to build a small database that holds the information about restriction enzymes contained in http://rebase.neb.com/rebase/link allenz ("allenz" standing for "all enzymes"). This version of the Rebase data provides substantially more information than the version we used in our earlier examples. We'll begin with the structure of the datafile, a program to read that data, and a design for the database based on it. We'll assume that the file is saved under the name allenz.data in the same directory that will contain the Python file with the code to process it.

The data

After some introductory text, which we'll skip as we've done in a number of other examples, the datafile contains an entry for each enzyme in the following format:

- <1> Enzyme Name
- <2> Prototype
- <3> Microorganism
- <4> Source
- <5> Recognition Sequence
- <6> Methylation Site
- <7> Commercial Availability
- <8> References
- ... blank line ...

The details of each field are as follows (conveniently, the numbers in angle brackets are actually part of the datafile format):

- 1. The *enzyme name* is the standard identifier of the enzyme (e.g., EcoRI).
- 2. The prototype is just the name of another of the file's enzymes, which was the first one discovered that recognizes the same sequence as the entry's enzyme. Entries for prototype enzymes have an empty prototype field.
- 3. The microorganism field consists of two or more subfields that name the genus, species, and possibly subspecies information. Some subspecies entries consist of somewhat informal multiword descriptions; others contain a single well-known subspecies designation.
- 4. The *source* is the name of an individual or National Culture Collection.
- 5. The recognition sequence is a coded description of both the sequence recognized by the enzyme and the way it cuts at that site.
- 6. The methylation site describes the location within the recognition sequence methylated by the cognate methylase; only about 10% of the enzymes in the file have information in this field. We will ignore this field.
- 7. Commercial availability is a series of single-letter codes indicating which vendors supply the enzyme. We will ignore this field.
- 8. As stated at the beginning of the datafile from which the rest of this information was extracted, "only the primary references for the isolation and/or purification of the restriction enzyme or methylase, the determination of the recognition sequence and cleavage site or the methylation specificity are given." The information in the references field is expressed as a list of one or more integers separated by a comma and a space. The integers refer to numbered references whose details are listed after the entry for the last enzyme.

The content of the recognition site field is a series of DNA bases, written for one strand only, from 5' to 3'. The cleavage site—if known—is represented by a caret (^), placed within the base sequence so that the cleavage occurs on the 3' side of the caret.

Some enzymes cleave several bases away from an end of the recognition site. For these, the base sequence in the field is followed by an indication of the location of the cut site(s). This is expressed with parentheses and slashes: for example, (5/10). The first number in this example indicates that the 5' to 3' cleavage occurs 5 bases downstream from the end of the recognition site, and although the complementary sequence is not shown, the second number indicates that the cleavage occurs 10 bases before the recognition site on the 5' to 3' complementary strand. Some enzymes cut on both sides of their recognition sequences, in which case the field will contain a parenthesized pair of numbers before and after the recognition sequence.

Reading the data

As we've seen in earlier examples, files often begin and/or end with information that is not part of the information we want to extract from them. The datafile we are working with in this case actually has two data sections, so the steps that skip, read, and stop at nondata must be performed twice. We'll read the data using an implementation of the Grand Unified Bioinformatics File Parser (see "The Grand Unified Bioinformatics File Parser" on page 146). The code is shown in Example 10-1.

```
Example 10-1. Reading Rebase "all enzyme" data
def read enzymes from file(filename):
    with open(filename) as file:
        skip intro(file)
        return (get enzymes(file), get references(file))
def skip intro(file):
    ""Skip through the documentation that appears at the beginning
    of file, leaving it positioned at the first line of the first enzyme""
    while not line.startswith('<REFERENCES>'):
        line = file.readline()
    while len(line) > 1:
                                                          # always 1 for '\n'
        line = file.readline()
    return line
def get enzymes(src):
    enzymes = \{\}
    enzyme = next enzyme(src)
    while enzyme:
        enzymes[enzyme[0]] = enzyme
                                                          # dict key is enzyme's name
        enzyme = next enzyme(src)
    return enzymes
def read field(file):
    return file.readline()[3:-1]
```

```
def read other fields(file):
    """The name of the enzyme has already been read;
    read the rest of the fields, returning a 7-tuple"""
    return [read field(file) for n in range(7)]
                                                             # read 7 fields
def next enzyme(file):
    """Read the data for the next enzyme, returning a list of the
    form: [enzyme_name, prototype, source, recognition_tuple,
    (genus, species, subspecies), references_tuple]"""
    name = read field(file)
    if name:
                                                             # otherwise last enzyme read
        fields = [name] + read other fields(file)
        fields[2] = parse organism(fields[2])
        fields[7] = [int(num) for num in fields[7].split(',')]
        file.readline()
                                                             # skip blank line
        return fields
def parse organism(org):
    """Parse the organism details"""
    parts = org.split(' ')
    if len(parts) == 2:
        parts.append(None)
    elif len(parts) > 3:
    parts[2:] = ' '.join(parts[2:])
    return tuple(parts)
def skip reference heading(file):
    """Skip lines until the first reference"""
    line = file.readline()
    while not line.startswith('References:'):
        line = file.readline()
    file.readline()
                                                             # skip following blank line
def next reference(file):
    """Return tuple (refnum, reftext) or, if no more, (None, None)"""
    line = file.readline()
    if len(line) < 2:</pre>
                                                             # end of file or blank line
        return (None, None)
    else:
        return (int(line[:4]), line[7:-1])
def get references(file):
    """Return a dictionary of (refnum, reftext) items from the
    references section of file"""
    # using a dictionary here because there are many references
    # and an enzyme may reference several of them
    refs = \{\}
    skip_reference_heading(file)
    refnum, ref = next_reference(file)
    while refnum:
        refs[refnum] = ref
        refnum, ref = next reference(file)
    return refs
```

```
if name == " main ":
    if len(sys.argv) < 2:</pre>
        filename = 'data/Rebase/allenz.data'
    elif len(sys.argv == 2):
        filename = sys.argv[1]
        print('Usage: read enzymes [filename]')
    enzymes, references = read enzymes from file(filename)
    print('Read', len(enzymes), 'enzymes and',
                                                        # print len instead of
          len(references), 'references')
                                                        # contents of huge lists
```

A schema for the Rebase database

There are many graphical conventions for representing database schema, but we don't need anything fancy. Figure 10-1 should suffice. What it shows is:

- The name of each table
- The names of the columns of each table
- A 1-N relationship from Enzyme to Organism, meaning there is only one Organism for each Enzyme, but there may be more than one Enzyme for each Organism
- A 1-N relationship from the Enzyme to itself, meaning there is (optionally) one prototype Enzyme for each Enzyme, but a single Enzyme can be the prototype of many others.
- A M-N relationship, represented in a separate table, between the Enzyme and Reference tables

The 1-N relationships are drawn the way they are because the relationship is represented in the database by a column in the N-side table that contains the key of the 1-side table.

Table 10-11 is a textual representation of the diagram in Figure 10-1.

Table 10-11. A textual schema for the Rebase data

Table	Column	Туре	Null?	Foreign key
Organism	*OrgID	integer	N	
	Genus	varchar(255)	N	
	Species	varchar(255)	N	
	Subspecies	varchar(255)		
Reference	*RefID	integer	N	
	Details	varchar(255)	N	
Enzyme	*Name	varchar(255)	N	
	Prototype	varchar(255)		Enzyme.Name ^a

Table	Column	Туре	Null?	Foreign key
	OrgID	integer	N	Organism.OrgID
	Source	varchar(255)	N	
	RecognitionSeq	varchar(255)	N	
	TopCutPos	integer		
	BottomCutPos	integer		
	TopCutPos2	integer		
	BottomCutPos2	integer		
EnzymeReference	*Enzyme	varchar(255)	N	Enzyme.Name
	*RefID	varchar(255)	N	Organism.OrgID

^a Enzyme. name is a key back to the same table, not a foreign key, but it serves the same purpose—the establishment of a 1-N relationship. There is nothing strange about one row in a table being related to another.

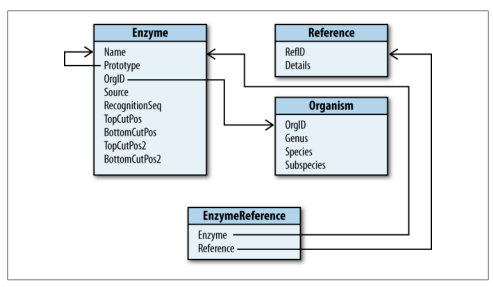


Figure 10-1. Schema diagram for the Rebase example

Using Relational Data

Given the preceding introduction and example, we can now look at the ways database tables and their contents are manipulated in practice.

SOL Basics

There are three major kinds of SQL operations:

Modify the schema

Add, remove, or modify tables.

Modify table contents

Add, remove, modify, or replace rows of tables.

Query the database

Get results matching database queries.

These operations are performed using SQL statements, as shown in Table 10-12.

Table 10-12. SQL statements

Category	Statement	Operation
Modify database schema	CREATE TABLE	Add table
	DROP TABLE	Remove table
	ALTER TABLE RENAME TO	Rename table
	ALTER TABLE ADD COLUMN	Add a column
Modify table contents	INSERT INTO	Add row(s)
	DELETE	Remove row(s)
	UPDATE	Modify a row(s)
	REPLACE	Replace a row(s)
Query database	SELECT	Obtain results

The rest of this chapter describes SQL fundamentals and how SQL is used from Python. As it proceeds, it uses an extended narrated example. Certainly much more could be done with this example, just as much more could be explained about SQL. The material here is intended to be consistent in depth with the material in the rest of this book.

Using the sqlite3 module

Large-scale database systems provide interactive interpreters, graphical user interfaces, and programs for executing SQL statements contained in text files. All of that is beyond the scope of this book and even of much of the work programmers do to use a database. What's important for programming purposes is that database systems provide libraries that implement functions (or methods, if object-oriented) for other code to call. We will use the SQLite3 database[†] and Python's sqlite3 module that interfaces to it because it is easy to use, it does not require a separate server, and it is included in the Python installation.

[†] You can find further information about SQLite3 at http://www.sqlite.org.



Python defines an abstract interface to SQL database systems. Accessing a specific database system requires a module that implements the interface for that kind of database. Implementations of the interface for many server-based systems can be found through the Python Package Index.‡

The sqlite3 module is an interface to a SQLite3 library compiled into the Python interpreter, so it doesn't require SQLite3 to be installed on your computer. However, if SQLite3 has been installed there will be a command-line interpreter program you can run called sqlite3.§

If you use the command-line sqlite3 program, bear in mind that standard interactive SQL tools require each statement to end with a semicolon.

While SQL is supposedly a standardized language, many database systems extend it in various ways and/or fail to implement some of its features. It is therefore always somewhat risky to show how to do things for relational databases in general, since a given SQL statement might not work with a specific database system. The SQLite3 database system is intentionally quite minimalist, but it nevertheless implements a large portion of SQL in a fairly standard way. Even though SQLite3 does not use a database server, it provides the same sort of interface used by full-featured client/server systems.



You'll want to learn more about databases and the sqlite3 module to accomplish your goals, and perhaps learn about other database systems as well. Beyond the pragmatics of actually using one, this section should give you a fairly clear picture of how relational databases manage and manipulate data. Given that foundation, it shouldn't be too difficult to learn about other uses and other database systems.

Connecting to the database

The first step in using the sqlite3 module is to create a database connection object:

conn = sqlite3.connect(dbname)

where dbname is the name of the file containing the database. The sqlite3 module provides convenient high-level methods for connection objects that hide some lower-level details that are part of interacting with a database connection. We'll look at three of them here:

[‡] Located at http://www.python.org/pypi; it is unclear how long it will be before each of these is updated to run with Python 3.

[§] If SQLite3 has not been installed on your machine and you want to try installing it, go to http: //www.sqlite.org/download.html. There are several versions available for download. The one you want (as of this writing) is sqlite3-3.6.14.bin.gz for Linux, sqlite3-3.6.14-osx-x86.bin.gz for Intel Macs running 10.5+, or salite-3_6_14.zip for Windows. The numbers after the first "3" will vary according to release.

connection.execute("a SQL statement"[, values])

Executes the SQL statement, replacing placeholders in the statement (discussed momentarily) with the corresponding element of *values* (a sequence or dictionary). Results (if any) are returned as an iterable that produces tuples containing the values for one result.

connection.executemany("a SQL statement", values-sequence)

Executes the SQL statement once for each sequence of values in valuessequence, replacing placeholders in the statement with the corresponding values, as with *connection*.execute. This is primarily used for insertions of multiple rows into tables.

connection.executescript("a SQL script")

Executes multiple SQL statements. This is a convenient way to create all the tables of a database in one method call, as you'll see later in Example 10-4. Note that each statement must end with a semicolon.

Programmatic SQL interfaces provide mechanisms for parameter substitution into statements. (This is one of those areas that is not as standardized as it should be.) The way it works in sqlite3 is very simple. The second argument to execute or executemany can be either a sequence or a dictionary. The placeholder used with a sequence is a question mark; when the query is executed, each question mark is replaced by the corresponding element of the sequence. There must be exactly as many elements in the sequence as there are question marks in the query. If the values are supplied in a dictionary, the query contains keys of the dictionary preceded by a colon, and the dictionary must contain every key used in the query (though it may contain others as well). We'll see examples of both mechanisms later.

The sqlite3 module defines an exception class, sqlite3. Error. You should always embed calls to execute inside try statements that catch this error and print out information about what has happened. After printing the error the exception clause should reraise the error with raise, since it won't normally be able to fix the problem.

Changes SQL statements make to a database are accumulated without actually being made to the underlying file. To move the changes into the file, it is necessary to commit them. This is the rough equivalent of closing an output file. The SQL COMMIT statement serves the same purpose in all standard SQL tools.

Connection objects provide the following methods for managing their state:

connection.commit()

Writes the accumulated changes to the database

connection.abort()

Abandons the accumulated changes without writing them to the database close()

Closes the connection without committing the changes

The accumulation of changes before a commit or rollback is called a transaction. Transactions are a critical part of programming large-scale databases. They prevent incomplete sets of changes from being made when an error occurs. For example, a banking system must ensure that money withdrawn from one account is deposited into another, so it uses a transaction to ensure that either both changes occur or neither does. Also, databases manage transactions so that one program's changes don't interleave with another's—the banking system shouldn't update the balance based on what was in the account when its transaction started, because that amount may have been changed by another transaction in the meantime.



A useful feature of methods of sqlite3 module classes is that they implicitly open a transaction whenever a SQL statement that changes the database schema is encountered. They also commit the implicit transaction when the next statement that does not change the schema is executed. This avoids schema changes and content changes interfering with each other.

Creating the database

SQL statements to create the four tables of the Rebase database look something like those shown in Example 10-2. (Many variations are possible even for a particular SQL implementation.) SQL keywords are usually shown in uppercase, so we'll follow that convention here.

Example 10-2. Database table creation statements for Rebase data

```
CREATE TABLE Organism(
    OrgID integer PRIMARY KEY,
    Genus text NOT NULL,
    Species text NOT NULL,
    Subspecies text
    );
CREATE TABLE Reference(
    RefID integer PRIMARY KEY,
    Details text NOT NULL
    );
CREATE TABLE Enzyme(
    Name text PRIMARY KEY.
    Prototype text,
    OrgID integer NOT NULL REFERENCES Organism(OrgID),
    Source text NOT NULL,
    RecogSeq text NOT NULL,
    TopCutPos integer.
    BottomCutPos integer,
    TopCutPos2 integer,
    BottomCutPos2 integer,
    );
```

```
CREATE TABLE EnzymeReference(
    Enzyme text NOT NULL REFERENCES Enzyme(Name),
    RefID integer NOT NULL REFERENCES Reference(RefID),
    PRIMARY KEY(Enzyme, RefID)
    );
```

The beginning of each of these statements names a table. Then, for each of the table's columns, the statement lists its name, type, and zero or more *constraints*. Constraints on the table itself follow the list of columns. Basic column constraints include:

NOT NULL

A value must be assigned to the column when a row is inserted or changed.

PRIMARY KEY

The column is the table's primary key and, implicitly, NOT NULL.

UNIQUE

No two rows in the table may have the same value for the constraint's column.

DEFAULT value

This specifies the value to give to a column in newly created rows when no value is specified for that column.

REFERENCES TableName(ColumnName, ...)

The column is a foreign key corresponding to the column(s) of the same name in the specified table.

Constraints that may appear on the table itself include:

```
PRIMARY KEY(ColumnName[, ...])
```

An alternate form of PRIMARY KEY constraint; if the primary key involves more than one column it must be specified as a constraint on the table.

```
FOREIGN KEY(ColumnName[, ...]) REFERENCES TableName(ColumnName, ...)
```

An alternate form of FOREIGN KEY constraint; if the foreign key involves more than one column it must be specified as a constraint on the table.

The next template shows the general form of the SQL statement for creating a table.

SQL

Creating a Table

The CREATE TABLE statement gives the name of the table to create, followed by a list of column specifications and other information. With brackets indicating optional components and ellipses repetition, the basic form of SQL's table creation statement is:

```
CREATE TABLE table_name(column_name column_constraint ...,
                       column name column constraint ...,
                       [table_constraint [, table_constraint
```

If you are redefining a table and reloading its data, you should drop the table first. The statements shown in Example 10-3 would normally precede the creation statements in Example 10-2, allowing changes to be made to a program's table creation statements before it is reexecuted.

Example 10-3. Dropping tables from the Rebase database

```
DROP TABLE IF EXISTS Organism
DROP TABLE IF EXISTS Reference
DROP TABLE IF EXISTS Enzyme
DROP TABLE IF EXISTS EnzymeReference
```

Dropping a table removes both the table's contents and its definition from the database. The only variation on the DROP TABLE statement is that the IF EXISTS portion may be omitted. An attempt made to drop a nonexistent table without IF EXISTS raises an exception.

SQL

Dropping a Table

The DROP TABLE statement removes a table from the database. Normally you would include IF EXISTS because an exception will be raised if you try to drop a table that is not in the database.

```
DROP TABLE [IF EXISTS] tablename
```

The sqlite3 Python code to execute the drop and create statements to define our example database is shown in Example 10-4. We use the nonstandard convenience method Connection.executescript to drop and create all the tables in one call, and a closing commit to tell the database to accept all the changes.

Example 10-4. sqlite3 code to create the Rebase database

```
import sqlite3
conn = sqlite3.connect(datafilename)
    conn.executescript('''
DROP TABLE IF EXISTS Organism
DROP TABLE IF EXISTS Reference
DROP TABLE IF EXISTS Enzyme
DROP TABLE IF EXISTS EnzymeReference
CREATE TABLE Organism(
    OrgID integer PRIMARY KEY,
    Genus text NOT NULL,
    Species text NOT NULL,
    Subspecies text
    );
```

```
CREATE TABLE Reference(
    RefID integer PRIMARY KEY,
    Details text NOT NULL
    );
CREATE TABLE Enzyme(
    Name text PRIMARY KEY.
    Prototype text,
    OrgID integer NOT NULL REFERENCES Organism(OrgID),
    Source text NOT NULL,
    RecogSeq text NOT NULL,
    TopCutPos integer,
    BottomCutPos integer,
    TopCutPos2 integer,
    BottomCutPos2 integer,
    );
CREATE TABLE EnzymeReference(
    Enzyme text NOT NULL REFERENCES Enzyme(Name),
    RefID integer NOT NULL REFERENCES Reference(RefID),
    PRIMARY KEY(Enzyme, RefID)
    );
''')
    except: sqlite3.OperationalError as err:
        print(err, file=sys.stderr)
        conn.rollback()
                                       # abort changes
        raise
conn.commit()
                                       # commit changes
```

Tables can also be renamed, as shown in the next template.

SQL

Renaming a Table

A table can be renamed as follows:

ALTER TABLE tablename RENAME TO new name

Internal database mechanisms depend on column names and types in a number of important ways that make changing the name, type, or constraints of a column problematic or even impossible. Database systems differ in the details. Other than renaming, the only kind of modification to a table's definition that SQLite3 supports is adding a column.

SQL

Adding a Column

To add a column to a SQLite3 database:

ALTER TABLE tablename ADD COLUMN columnspec

The column specification is the same as in a CREATE TABLE statement, except that certain kinds of constraints may not be included.

Loading data into tables

The statement used to add a row to a table is INSERT INTO. Only a few of the insertion statements for our example database are shown, because altogether it will have approximately 15,000 rows!

SQL

Inserting a Row into a Table

The basic SQL statement for inserting a row into a table has the form:

```
INSERT INTO tablename VALUES(value1, value2, ..., valuen)
```

where n must be the number of columns in the table. The values must appear in the order of the corresponding columns.

When a program is doing insertions, a better approach is to use a question mark in place of each value and supply the actual values as the statement is repetitively executed:

```
connection.execute("INSERT INTO tablename
                   VALUES(?,?,?,...)",
                   value1, value2, ..., valuen)
```

As always, a dictionary may be used to supply the values instead of a sequence, with names preceded by colons in place of the question marks.

The first step in loading the data into the database is to read all the information from the datafile, as shown in Example 10-1. What that produces is a dictionary of enzyme information keyed by the enzyme names, and a dictionary of references keyed by the numbers as they appear in the datafile. The Reference table can be loaded directly from the references dictionary, but the other three tables require some manipulation of the raw data: we have to split the enzyme data into the information that goes into the Organism table and the information that goes into the Enzyme table, and we need to create the M-N table that connects the Enzyme and Reference tables.

Example 10-5 shows the code for loading the data from the text file into the database. The top-level function is **load data**. It calls a separate function for each table:

- load reference data
- load organism data
- load enzyme data
- load enzyme reference data

In addition, load enzyme data calls recognition info to parse the cut site string into its components. We're ignoring that detail here but leaving the function in as a "stub" that just returns a list of its sequence argument and four zeros. We will deal with these details at a later point.

```
Example 10-5. Loading the Rebase database
```

```
def load data(dbname, enzymes, references):
    """Reorganize the information in the enzymes and references dictionaries in accordance
    with the dbname's schema and load the data into the database's tables"""
    try:
        conn = get connection(dbname)
        load reference data(conn, references)
        organism ids = load organism data(conn, enzymes)
        load enzyme data(conn, enzymes, organism ids)
        load enzyme reference data(conn, enzymes)
        conn.commit()
    except sqlite3.OperationalError as ex:
        print(ex, file=sys.stderr)
        raise
                           # reraise exception so Python can handle it
def load reference data(conn, references):
    for refid, ref in references.items():
        store data(conn, 'Reference', (refid, ref))
def load organism data(conn, enzyme data):
    """Return a 'reverse' dictionary keyed by the tuples in the list enzyme_data, which have
    the form (Genus, Species, Subspecies); the dictionary's values are sequentially generated
    integer IDs, which will be used as foreign keys in the Enzyme table"""
    organism ids = {}
    for orgid, data in enumerate(enzyme data.values()):
        # generating OrgIDs as we go
        org = data[2]
        # this is the only part of an enzyme's data that is relevant here
        if not org in organism ids:
             store_data(conn, 'Organism', (orgid+1,) + org)
            organism ids[org] = orgid+1
    return organism ids
def load enzyme data(conn, enzymes, organism ids):
    for data in enzymes.values():
        store data(conn,
                     'Enzyme',
                    (data[0],
                                                         # name
                     data[1] or None,
                                                         # prototype (None if ")
                     organism_ids[data[2]],
                                                     # organism ID
                     data[3]) +
                                                         # source
                                                     # recognition sequence
                    recognition info(data[4]) +
                                                         # methylation site
def load enzyme reference data(conn, enzymes):
    """For each reference of each element of enzymes add an entry to the M-N EnzymeReference
    table with the name of the enzyme and the ID of the reference"""
    for data in enzymes.values():
```

```
for refid in data[7]:
                                                            # refid list
             store data(conn, 'EnzymeReference', (data[0], refid))
# temporary implementation: does nothing
def recognition info(seqdata):
    """Parse the recognition sequence data seqdata, returning a tuple of the form:
         (sequence, top_cut_pos, bottom_cut_pos, top_cut_pos2, bottom_cut_pos2)"""
    return (seq, 0, 0, 0, 0)
```

These four functions make use of the code shown in Example 10-6 to avoid lengthy program text and code repetition. The function make insert string generates an INSERT statement with the right number of question marks for a table of a given name and number of columns. STORE STMTS is a dictionary keyed by table name that contains the result of make insert string for each of the four tables of the database. The function store data encapsulates the execution of an INSERT statement inside a try statement. It obtains the appropriate string for the INSERT statement from the STORE STMTS dictionary.

Example 10-6. Utilities to support Rebase database loading

```
def make insert string(tablename, n):
    """Return an INSERT statement for tablename with n columns"""
   return ('INSERT INTO ' + tablename + ' VALUES ' +
            '(' + ', '.join('?' * n) + ')') # n comma-separated ?s
STORE STMTS = {tablename: make insert string(tablename, ncols)
               for tablename, ncols in
               (('Organism', 4),
                ('Reference', 2),
                ('Enzyme', 9),
                ('EnzymeReference', 2))}
def store data(conn, tablename, data):
    """Store data into tablename using connection conn"""
        conn.execute(STORE STMTS[tablename], data)
   except Exception as ex:
        print(ex)
        raise
```

SOL Queries

With tables defined and data loaded into the database, it is time to consider the crux of SQL use: queries that fetch information from the database. This final part of the chapter explores the key aspects of SQL data access. The topic is far too complex for us to cover anything but the most basic techniques, but the ones we will cover account for a significant proportion of what even a moderately experienced SOL programmer would use in practice. Also keep in mind that implementations in other databases may vary in some minor ways from the SQLite3 syntax described here.

Simple database queries

Creating and loading tables is essentially just administrative work. The real power and complexity—of SQL comes into play in accessing the information that's already stored in a database. A key advantage SQL has over directly accessing a program's data structures is that it provides many powerful mechanisms for choosing columns, combining data from multiple tables, imposing filters, performing calculations, and constructing temporary tables that can be used as the source for further accesses.

The heart of SQL is its SELECT statement. A SELECT statement specifies combinations of column names, table names, calculations, conditions, and so on used to refer to data. The execution of a SELECT statement produces a (possibly empty) stream of results from which the program can read.

SQL

Simple SELECT Statement

The simplest form of the SELECT statement is:

```
SELECT column name, ... FROM table name
```

The result is a stream of tuples containing values in the designated columns for every row of the specified table. To access all the columns of a table, use the shorthand:

SELECT * FROM table name

In documentation and discussion of SELECT statements, each part after the column names—FROM table name, for instance—is referred to as a clause. SELECT statements can have a variety of optional clauses. An important one in practice is LIMIT, since for tables with many rows, such as those in our Rebase example, you probably won't want to access all of them.

SQL

Limiting SELECT Statement Results

You can limit the number of rows a query returns as follows:

```
SELECT column name, ... FROM table_name LIMIT number
```

You can also add another clause to SELECT statements that have a LIMIT clause to say where in the sequence of results you want to start (the "offset"):

```
SELECT column name, ... FROM table name LIMIT number OFFSET position
```

Another option is to add a clause specifying the column(s) to use in determining the order of the results:

SELECT column name, ... FROM table name ORDER BY column name

Column names are not the only thing that can follow the SELECT keyword. A wide range of expression types can be used, with syntax much like Python's. (An important exception is that a single equals sign is an equality test; there is no assignment.)

SQL

Expressions in SELECT Statements

Some of the more important expressions are:

```
column name
literal_value
expression(s)
expression1 binary operator expression2
expresssion IS/NOT NULL
expression1 [NOT] LIKE expression2
expression1 [NOT] BETWEEN expression2 AND expression3
function name(expression, ...)
```

("Literal" values are numbers, single-quoted strings, NULL, etc.) SQL functions you can use in SELECT statements include COUNT, MAX, MIN, AVG, and SUM. An expression following LIKE includes wildcard characters, with matching a single character and % zero or more. Matching is case-insensitive.

Another maneuver that is often important is to limit expressions—COUNT(), in particular—to distinct values. This is like the difference between a set and a tuple, in that sets do not contain duplicate values.

SQL

DISTINCT in SELECT Statements

The SQL keyword DISTINCT can appear before the expressions of a SELECT statement or the arguments to a function (inside the parentheses). This qualifies the query so that only one row for each distinct value is included in the results.

```
SELECT COUNT(DISTINCT Genus, Species) FROM Organism
```

Example 10-7 shows some simple SELECT statements for the Rebase database. Example 10-8 shows code to execute them in Python. The code includes a convenience function to write some output since conn.execute() just returns an iterable object—it doesn't print anything.

```
Example 10-7. Simple SELECT statements in SQL
```

```
SELECT Name, Prototype FROM Enzyme LIMIT 5
SELECT * FROM Organism LIMIT 4 OFFSET 6
SELECT * FROM Organism
        ORDER BY Genus, Species, Subspecies
```

```
LIMIT 4 OFFSET 6
SELECT COUNT(*) FROM Organism
Example 10-8. Simple SELECT statements using the sqlite3 module
dbname = 'path to database'
conn = sqlite3.connect(dbname)
def execute(query, args=[]):
   print()
   print(query)
   print()
   try:
        for result in conn.execute(query, args):
            print(result)
   except sqlite3.Error as ex:
        print(ex, file=sys.stderr)
for query in (
    'SELECT Name, Prototype FROM Enzyme LIMIT 5',
    'SELECT * FROM Organism LIMIT 4 OFFSET 6',
    '''SELECT * FROM Organism
           ORDER BY Genus, Species, Subspecies
           LIMIT 4 OFFSET 6''',
    'SELECT COUNT(*) FROM Organism',
    'SELECT MAX(Name, FROM Enzyme',
    'SELECT COUNT(Subspecies) FROM Organism',
    'SELECT COUNT(DISTINCT Genus) FROM Organism',
    '''SELECT DISTINCT Species FROM Organism
              WHERE Genus = 'Mycobacterium'
              ORDER BY Species, Subspecies ''',
   ):
   execute(query)
conn.close()
The results of executing the preceding script are shown next. The third query reorders
the results first by genus, then species, and finally subspecies:
    SELECT Name, Prototype FROM Enzyme LIMIT 5
    ('HpyF30I', 'TaqI')
    ('TmiI', '')
    ('BstD102I', 'BsrBI')
    ('BssHII', 'BsePI')
('BstXII', 'MboI')
    SELECT * FROM Organism LIMIT 4 OFFSET 6
    (7, 'Bacillus', 'stearothermophilus', '1473')
    (8, 'Actinobacillus', 'suis', 'NH')
(9, 'Helicobacter', 'pylori', 'RFL21')
```

(10, 'Thermus', 'species', None)



That the results of the second query are ordered according to the numerical IDs we generated for them is an implementation detail that you can't generally rely on.

```
SELECT * FROM Organism
               ORDER BY Genus, Species, Subspecies
               LIMIT 4 OFFSET 6
(2400, 'Acetobacter', 'pasteurianus', None)
(1259, 'Acetobacter', 'pasteurianus', 'B')
(1538, 'Acetobacter', 'pasteurianus', 'C')
(1899, 'Acetobacter', 'pasteurianus', 'D')
SELECT COUNT(*) FROM Organism
(3031,) # note that even single values are returned as tuples
SELECT MAX(Name) FROM Enzyme
('Zsp2I',)
SELECT COUNT(Subspecies) FROM Organism
           # the number of organisms whose subspecies is not NULL
(2530,)
SELECT COUNT(DISTINCT Genus) FROM Organism
(241,)
```



There are ordering dependencies among the kinds of clauses that can appear in a sqlite3 SELECT statement. | Essentially, whatever clauses are contained in query must appear in the following order:

```
DISTINCT
EXPRESSION(s)
FROM
WHERE expression
GROUP BY column names
ORDER BY column names
LIMIT integer [OFFSET integer]
```

The definition of execute in Example 10-8 takes an optional second argument. This is for the values to be substituted into a parameterized query. Here are two of the queries we just showed, but in parameterized form. We use tuples to supply the query and the

A diagram documenting all the variations on SELECT together with explanatory text is available at http://www.sqlite.org/lang_select.html. Because this is complete documentation much of it is more advanced than you will want to deal with, but you should be able to pick out the important parts.

values to be substituted. The first query uses a question mark as a placeholder, while the second uses the dictionary notation:

```
for query in (
    ("SELECT Name, Prototype FROM Enzyme LIMIT ?", (5,)),
    ("SELECT * FROM Organism LIMIT :lim OFFSET :off", {off: 6, lim: 5})
    execute(query[0], query[1])
```

Qualified database gueries

Despite the variety of queries we've seen so far, the most significant part of a SELECT statement is the ability to restrict results based on column values. This is done using WHERE clauses and expressions. WHERE clauses follow the FROM clause(s) in SELECT statements.

SQL

WHERE Clauses in SELECT Statements

A basic WHERE clause has the form:

```
SELECT * FROM table name WHERE expression
```

Often, the expression has multiple subexpressions connected by combinations of AND and OR:

SELECT * FROM tablename WHERE expression₁ AND expression₂

Example 10-9 shows two queries with WHERE clauses.

```
Example 10-9. WHERE clauses in SELECT statements
for query in (
    '''SELECT COUNT(*) FROM Enzyme WHERE Prototype IS NULL''',
    '''SELECT DISTINCT Species FROM Organism
              WHERE Genus = 'Mycobacterium'
              ORDER BY Species''',
    ):
    execute(query)
Here are the results:
    SELECT COUNT(*) FROM Enzyme WHERE Prototype IS NULL
    (606,)
    SELECT DISTINCT Species FROM Organism
                  WHERE Genus = 'Mycobacterium'
                  ORDER BY Species
    ('avium',)
    ('butyricum',)
```

```
('chelonei',)
('fortuitum',)
('gordonae',)
('habana',)
```

Relationship queries

Consider these example queries performed in the sqlite3 interactive command shell:

SELECT * FROM 0	rganism WHERE OrgID H	BETWEEN 11 AND 14;	
OrgID	Genus	Species	Subspecies
11	Citrobacter	freundii	RFL22
12	Streptomyces	luteoreticuli	
13	Streptomyces	albohelvatus	
14	Mycobacterium	butyricum	
SELECT Name, Or	gID FROM Enzyme WHER	OrgID BETWEEN 11 AND	14;
Name	OrgID		
Cfr22I	11		
C†r22I SluI	11 12		
			
SluI	12		
SluI SabI	12 13		
SluI SabI M.MbuII	12 13 14		

The results of these queries show that rows in the Enzyme table with an OrgID value of 14 represent the enzymes produced by the organism whose OrgID is 14. Suppose we wanted to start with the genus Mycobacterium, species butyricum, and subspecies NULL and find the enzymes that organism produces. First we can find the OrgID of the row representing the organism:

```
SELECT * FROM Organism
        WHERE Genus = Mycobacterium AND
        Species = butyricum AND
        Subspecies IS NULL;
OrgID
                                                         Subspecies
                   Genus
                                      Species
                  Mycobacterium
                                      butyricum
```

Now that we have the organism's ID, we can use it in a query to get the enzymes it produces:

```
SELECT Name FROM Organism WHERE OrgID = 14
Name
M.MbuII
M.MbuIV
M.MbuT
M.MbuIII
```

We can write a function that generalizes these steps to return a list of the enzymes any organism produces, but it must handle an awkward problem. When using question marks in queries, the second argument to execute must be a sequence with as many elements as there are question marks in the query. However, to match a row where a column value is NULL requires the WHERE clause to include IS NULL for the column, rather

than = ?. Therefore, the function must prepare two different queries according to whether or not a value is supplied for the subspecies:#

```
def get organism enzymes(genus, species, subspecies=None):
    query = ('''SELECT OrgID from Organism
                WHERE genus = ? AND species = ? AND subspecies ''' +
             (' = ?' if subspecies else 'IS NULL'))
    rows = conn.execute(query,
                         (genus, species, subspecies) if subspecies
                         else (genus, species))
    if rows:
        lst = list(rows)
                                                       # a tuple of tuples
        assert len(lst) == 1, len(lst)
        orgid tuple = lst[0]
                                                       # the tuple (orgid, ) retrieved by the
                                                       # first query
        assert len(orgid tuple) == 1, len(orgid tuple)
    return [row[0] for row in
            conn.execute('''SELECT name FROM Enzyme WHERE OrgID = ?''',
                          orgid tuple)]
```

The first query gets the OrgID for the organism with the specified genus, species, and subspecies. The second query uses that ID to retrieve from the Enzyme table all enzymes with an OrgID that matches the one the first query retrieved. This corresponds to the interactive queries shown previously.

These examples demonstrate how 1-N relationships work. A primary key in one table is a foreign key in another table. While primary keys must be unique among the rows of a table, foreign keys are not restricted in that way. The rows with a particular value for their foreign key are the N rows of the 1-N relationship, with that value selecting a unique row of the other table. (There could be several foreign keys in one table; we're talking about just one here for simplicity's sake.) Figure 10-2 illustrates the process.

This kind of two-step lookup for 1-N relationships is so common that SQL provides syntax expressing it in a single query. The definition of get organism enzymes shown earlier can be greatly simplified by using this syntax:

```
def get organism enzymes query(genus, species, subspecies=None):
    query = ('''SELECT Enzyme.name from Organism, Enzyme
                  WHERE genus = :genus AND
                         species = :species AND
                         subspecies ''' +
               (' = :subspecies' if subspecies else 'IS NULL') +
'''AND Organism.OrgID = Enzyme.OrgID''')
    return [row[0] for row in
             conn.execute(query,
                            {'genus': genus,
                             'species': species,
                             'subspecies': subspecies}
                            )]
```

#It isn't the question marks that are the problem—even if the function were to construct a literal query with no question marks, either by string concatenation or the string formatting mechanism, it would still have to construct a different query for when the subspecies is None and for when it's not.

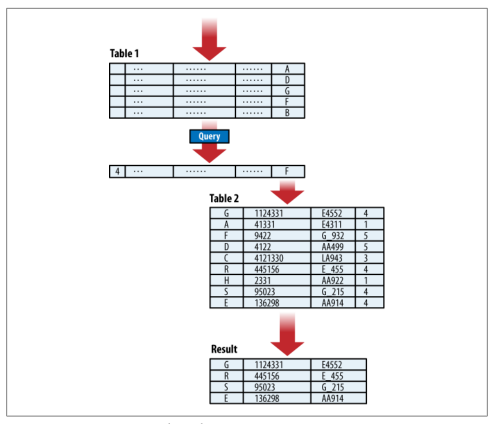


Figure 10-2. Navigating a 1-N relationship

This is the first query we've shown that accesses more than one table. Two tables may have a column with the same name, so to disambiguate which table's column is meant the name of the table and a period are added to the column name. (This is analogous to how names defined in a Python module are accessed by prefixing them with the module's name and a period.)



Even if a column name is unique among the tables of the query, qualifying each column name with the name of its table can help make the query clearer.

The critical piece of the new query is the first line that's highlighted. It restricts the results to just those rows of the Enzyme table whose foreign key OrgID matches the OrgID of the row of the Organism table with the values of genus, species, and subspecies.

TEMPLATE

Querying 1-N Relationships

Given the following:

- *key* is the primary key of *table1*.
- foreignkey is the foreign key from table2 to table1.
- *criterial* is a part of a SELECT statement that obtains a single row from *table1*.
- resultcolumns2 are the columns of table2 to return as results.

queries to obtain those results take the following form:

```
SELECT resultcolumns2
      FROM table1, table2
      WHERE criteria1 AND
             tablename1.key = tablename2.foreignkey
```

The syntax shows only single-column keys, but multicolumn keys can be used by adding more AND phrases to the WHERE clause.

Working with M-N relationship tables is actually no different than working with the foreign keys from one table to another that implement 1-N relationships. In most respects an M-N relationship table just implements two separate 1-N relationships, one with each of the two tables it connects. Each of the two 1-N relationships uses a different foreign key. M-N relationships are really a design concept; the database mechanisms used to implement them are the same.

One difference does show up in queries involving M-N tables, though. We saw earlier how getting the "N" side of a 1-N relationship involves first finding the "1" row according to specified criteria, then getting each row from the "N" side whose foreign key corresponds to the primary key of the row on the "1" side. If we want to go from a row in one table to all the rows in another table by way of an M-N table, we have three tables to consider. Figure 10-3 illustrates the process.

Our Rebase database contains an M-N table that indicates what references apply to which enzymes. This must be an M-N table because an enzyme can have multiple references, and one reference can be for multiple enzymes. A typical query, though, would ask either for the references corresponding to a specific enzyme or for the enzymes corresponding to a specific reference:

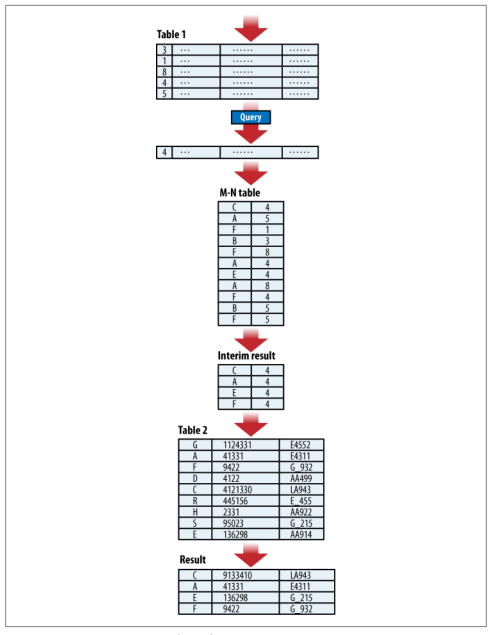


Figure 10-3. Navigating an M-N relationship

```
def get_references_for_enzyme(enzyme):
     ""Return the reference details for the enzyme named enzyme"""
    return [row[0] for row in
            conn.execute(''
                SELECT Reference.details
                FROM Enzyme, Reference, EnzymeReference
                WHERE Enzyme.name = ? AND
                       Enzyme.name = EnzymeReference.Enzyme AND
                       Reference.refID = EnzymeReference.RefID
                       ''', (enzyme,))]
def get enzymes for reference(ref):
    """Return the name of the enzymes referenced
    by the reference whose ID is ref"""
    return [row[0] for row in
            conn.execute('
                SELECT Enzyme.name
                FROM Enzyme, Reference, EnzymeReference
                WHERE Reference.RefID = ? AND
                       Enzyme.name = EnzymeReference.Enzyme AND
                       Reference.refID = EnzymeReference.RefID
                       ''', (ref,))]
```



In either case, the "specific" item is obtained through the usual selection criteria, but its key column(s) match multiple rows of the M-N table rather than of another table from a 1-N relationship. Then, the foreign keys to the other side of the relationship are used to access rows in the other table.

The following template sums up the details of queries for M-N relationships.

TEMPLATE

Querying M-N Relationships

Given the following:

- *key1* is the primary key of *table1*.
- *key2* is the primary key of *table2*.
- table3 represents an M-N relationship between table1 and table2.
- foreignkey1 is the foreign key from table3 to table1.
- foreignkey2 is the foreign key from table3 to table2.
- *criterial* is a part of a SELECT statement that obtains a single row from *table1*.
- resultcolumns2 are the columns of table2 to return as results.

the form of the query to get those results is as follows:

```
SELECT resultcolumns2
      FROM tablename1, tablename2, tablename3
      WHERE criteria1 AND
```

```
tablename1.key1 = tablename3.foreignkey1 AND
tablename2.key2 = tablename3.foreiqnkey2
```

The syntax shows only single-column keys, but multicolumn keys can be used by adding more AND phrases to the WHERE clause.

Querying the Database from a Web Page

In this final section of the chapter we'll put a web-based user interface in front of the database. There are a number of strategies for accomplishing this based on HTML forms. These include:

- Allowing the user to type an arbitrary query into a text box*
- JavaScript-driven dynamic pages that change what's presented in the form according to the information already entered, the way some widely used web search pages
- Elaborate "query by example" dynamic web pages
- A fixed form that provides fields for specifying values for a particular query
- A form that presents a menu of guery forms
- A page that presents multiple forms, each with fields for specifying values for a particular query

In order to keep the HTML details as simple as possible, we'll use the last approach.

HTML frames

We are going to use HTML frames to construct a page with a choice of queries on top and results on the bottom. Basic HTML frame arrangements are actually quite simple: the frame page just names the other pages and says how much horizontal or vertical space each should take. The amount of space is expressed either in pixels, as percentages of the parent page, or with an asterisk (meaning that it should take all the remaining space). Example 10-10 shows the HTML for our frame page.

Example 10-10. HTML page for Rebase browser-based queries

```
<html>
<head>
<title>Rebase Database Oueries</title>
</head
 <frameset rows="400,*">
   <frame name='queries' src='allenz.html' scrolling='yes'>
   <frame name='results' scrolling='yes'>
```

* The obvious security problem of this approach can easily be obviated by requiring the first word of the query to be SELECT. The real problem with this approach is the requirement that the user know enough SQL to formulate the desired queries, as well as the schema of the target database.

```
</frameset>
</html>
```

Note that this page does not include a body—a frameset is used instead. The rows attribute in the <frameset> tag specifies that the top frame should be 400 pixels high, and the bottom frame should occupy the rest of the window. Each of the two <frame> tags specifies a name and says to activate scrollbars when needed. The src attribute of the first <frame> tag gives the address (relative, in this case) of the web page to be displayed in that frame. The second frame starts out empty: it will be filled with the results of the queries performed in the first frame.

An advantage of this two-frame approach is that the contents of the top frame are stable, making it easier for a user to find queries within it. Another advantage is that the form does not have to be sent back to the browser with each query's results. † The results will replace the contents of the bottom frame, leaving the top frame unaffected.

An HTML page of guery forms

The HTML page in the top frame will contain little more than a series of forms. We'll avoid discussing HTML tables and Cascading Style Sheets (CSS) and consider only the simplest possible construction (however, you can find several variations for this page that use tables, CSS, and even JavaScript in the code files accompanying the book). Each query will be displayed on the web page using HTML like that in Example 10-11.

Example 10-11. Simple HTML for query forms

```
<form action='/cgi-bin/rebase' target='results'>
    <input type='hidden' name='query' value='species for genus'>
     genus <input name='genus'> (e.g. Vibrio)<br/>
     <input type='submit'>
    </form>
```

This will display in a browser something like what's shown in Figure 10-4. Here's a line-by-line description of what the HTML means:

- **1** The title of the query, in bold.
- **2** The beginning of a form (on a new line).
 - a. The form's action is the program *rebase* found in the *cgi-bin* subdirectory of the directory from which the server is running.[‡]

[†] This was discussed at the end of the previous chapter; see Example 9-20 ("HTML forms with CGI scripts" on page 350).

[‡] See "Serving CGI requests" on page 343 for instructions on how to run a simple CGI server using Python's http.server module.

- b. The form specifies the target 'results', which is the name of the other frame; if no target were specified the results of the query would replace the page with the form in the browser.
- An input field with type 'hidden'; this kind of field is not displayed but is included with the rest of the fields when the form is submitted—in effect, it is an invisible constant field. It is this field that tells the Python program which SQL query to execute.
- A label, a one-line text input field, and an example to show the user the sort of value the query expects the field to contain.
- The submit button; when this is pressed, the form's action is submitted to the web server along with the field values. With no 'value' attribute the button will have a default label, which for most browsers will be "Submit."
- **6** The end of the form.

Number of Species b	y Genus
minimum count	(e.g. 3)
Submit	

Figure 10-4. Display of a simple query form

Multiple forms can appear on the same page, as Example 10-12 illustrates. Pressing a submit button invokes the action of the form that contains it with the values of that form's fields. There is no connection to the fields of other forms on the same page.

Example 10-12. A page of Rebase queries

```
<html>
<head>
<title>Rebase Oueries</title>
</head>
<body>
<b>Number of Species by Genus</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='number of species by genus'>
   minimum count <input name='minimum count'> (e.g. 3) <br/>
    <input type='submit'>
 </form>
<br/>
<br/>
b>Number of Enzymes by Organism</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='number of enzymes by organism'>
   minimum count <input name='minimum count'> (e.g. 3) <br/>
    <input type='submit'>
 </form>
<b>Species for Genus</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='species for genus'>
```

```
genus <input name='genus'> (e.g. Vibrio)<br/>
    <input type='submit'>
 </form>
<br/>
<br/>
Subpecies for Genus and Species</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='subspecies_for_genus_species'>
   genus <input name='genus'> (e.g. Thermus)<br/>>
   species <input name='species'> (e.g. thermophilus)
   <input type='submit'>
 </form>
<br/>tb>Enzymes for Organism</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='enzymes for organism'>
   genus <input name='genus'> (e.g. Helicobacter)<br/>>
   species <input name='species'> (e.g. pylori)<br/>
    (optional) subspecies <input name='subspecies'> (e.g. RFL44)
    <input type='submit'>
 </form>
<b>Prototype Enzymes</b>
 <form action='cgi-bin/rebase' target='results'>
    <input type='hidden' name='query' value='prototype_enzymes_like'>
   name like <input name='name like'> (e.g. Ha%)<br/>>
    <input type='submit'>
  </form>
<br/>
<br/>
<br/>
d>References for Enzyme</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='references_for_enzyme'>
   enzyme <input name='enzyme'> (e.g. EcoRI)<br/>>
    <input type='submit'>
 </form>
<br/>
<br/>
b>Enzymes for Reference Number</b>
 <form action='cgi-bin/rebase' target='results'>
 <input type='hidden' name='query' value='enzymes for reference'>
   reference number <input name='number'> (e.g. 11)
    <input type='submit'>
 </form>
</body>
</html>
```

This page by itself will display as in Figure 10-5. The frame-based web page is shown in Figure 10-6.

Tips, Traps, and Tracebacks

Tips

 Just as you should develop and debug regular expressions using a testing tool like re-try, it is often helpful to use a tool to explore a database directly. If your computer has the command-line command sqlite3 installed, you could use that. (If it doesn't, you can try downloading the appropriate file for your platform from

Number of Species	by Genus
minimum count	(e.g. 3)
Submit	
Number of Enzyme	es by Organism
minimum count	(e.g. 3)
Submit	
Species for Genus	
genus	(e.g. Vibrio)
Submit	
Subpecies for Genu	
genus	(e.g. Thermus)
species	(e.g. thermophilus)
Enzymes for Organ genus species	(e.g. Helicobacter) (e.g. pylori)
(optional) subspecies	
Submit Prototype Enzymes	
name like	(e.g. Ha%)
Submit	
References for Enz	yme
enzyme	(e.g. EcoRI)
Submit	
Enzymes for Refere	ence Number
reference number	(e.g. 11)
Submit	

Figure 10-5. Display of a page of query forms

http://www.sqlite.org/download.html—you want just the file that contains the precompiled command-line installation, or, if there is none for your platform, the source code.) There are many GUI interfaces—some free, some commercial; some platform-independent, some platform-specific. You can find a long list of information about them at http://www.sqlite.org/cvstrac/wiki?p=ManagementTools. A particularly convenient, simple, free, and cross-platform GUI interface to

Number of Specie	es by Genus
minimum count	(e.g. 3)
Submit	
Number of Enzyn	mes by Organism
minimum count	(e.g. 3)
Submit	
Species for Genus	IS
genus Vibrio	(e.g. Vibrio)
Submit	
Subpecies for Ger	enus and Species
genus	(e.g. Thermus)
species	(e.g. thermophilus)
Submit	500 8600 8 86 00
species for genus	s, genus=Vibrio
anguillarum	
cholerae	
fischerii	
harveyi	
nereis	
nigripulchritu	
parahaemoly	yticus
species	
vulnificus	

Figure 10-6. Frame-based display of a page of query forms

sqlite3 is the SQLite Database Browser, which you can download from http:// sqlitebrowser.sourceforge.net/index.html.

• You can save yourself a lot of trouble by always using single-column integer-valued primary keys in relational database tables. This chapter's examples used some other kinds in order to demonstrate how keys work, but in practice there is a major problem with using meaningful column values as keys: you may need to change those values while curating the database (e.g., to correct inconsistent abbreviations, spelling, or names), but database systems generally do not allow changes to the values in primary key columns. Even if the database system did allow changes to foreign key values, making them would require the same changes to be made in all the rows of any other tables that use the key values as foreign keys. Using integervalued primary keys—preferably ones that are automatically generated by the database system—avoids these problems.

- The implementation of SQLite3 makes using an integer as a primary key easy, because every table has a key called ROWID that is automatically incremented every time a row is inserted into the table.
- You can simplify calls to Connection.execute by naming query strings elsewhere and just using their names when calling execute. This also has the advantage that the same query string can be used in multiple places.

Traps

- sqlite3.Connection.close does not do a commit; you must call commit before closing the connection if you want the changes to be committed.
- If you are working in the interpreter and don't close a sqlite3 database connection, attempting to open it again will raise an error. This can happen very easily when you are debugging your code. You should always close connections in the finally of a try statement.

Tracebacks

Error messages you might encounter while using the sqlite3 module include the following:

sqlite3.Error

An error has been produced by the sqlite3 module.

sqlite3.OperationalError

An attempt has been made to execute an invalid SQL statement.

SyntaxError: invalid syntax

A string passed to eval contained a syntax error. (Python must parse the string before it can execute it.)

Structured Graphics

Bioinformatics deals with enormous amounts of data. People process visual imagery far more easily and quickly than words and numbers, so many bioinformatics websites and applications provide highly effective visualizations of complex data. Generating graphical presentations is an important aspect of bioinformatics programming. This chapter will show you how to do it.

We will look at implementations of three kinds of information displays:

- *Histograms* are useful for anything involving counts. They are used widely in many fields and business operations. In bioinformatics, they are frequently used to provide high-level visual overviews of quantitative distributions.
- Dot plots are also a general-purpose chart type, but in bioinformatics they have special uses for visualizing how two sequences—or a sequence and itself—are related.
- *Raw data* collected from laboratory devices such as sequencing machines is usually displayed as curves. The classic example is the four-colored "trace" of the data from a traditional sequencing device.

The data used for all of the examples can be found on the book's website. In writing your own applications, you can obtain similar data via URL queries.

Introduction to Graphics Programming

Computers represent visual images in one of two ways:

Bitmaps

A bitmap is a two-dimensional arrangement of *pixels*, each representing the color of a "dot" at a particular location in the image.

Structured graphics

A structured image is a set of instructions in a "drawing language" that describe individual objects representing different kinds of shapes and text.

You may already be familiar with this distinction from having used some applications that manipulate bitmaps and others that manipulate diagrams. Photographs, icons, screen snapshots, and the contents of the display itself are represented as bitmaps. Images in which you can select individual components, such as a rectangle or piece of text, are structured. You can add, remove, and group components in a structured image, and change a given component's properties (color, location, border width, etc.).

A structured image may be converted to a bitmap, as when a snapshot is taken of a window on the screen displaying a diagram. Bitmaps, though, cannot be turned into structured representations, because they do not contain any information about the individual components of the image.

Concepts

It is difficult to talk about graphics programming in the abstract. Structured graphics are implemented in a variety of ways, using numerous file formats, languages, libraries, and tools. Some technologies store images using file formats resembling the object persistence mechanisms discussed at the end of Chapter 6, while others represent them in an XML-based format. Still others are designed to display images in windows on the screen.

For the most part, all graphics programming technologies share a common set of concepts and mechanisms. However, the vocabulary and technical details used in working with these concepts and mechanisms can vary significantly from one technology to another. Learning to program in your first graphics technology can be quite disconcerting because you must learn not only a new conceptual framework and new terminology, but also all the details of that particular technology.

Structured graphics can be discussed and even programmed without anything more than a vague idea of what output device will be used to display them. In fact, a diagram or image can be generated without any display at all. The result is either a file written in some structured graphics format, a textual description written in a well-defined graphics language, or simply a data structure inside the computer's memory. Outputting the graphics to a screen, printer, or other device is essentially an independent step. The three main aspects of all structured graphics tools are their coordinate system, components, and component properties.

Coordinate system

Structured graphics are created within a coordinate system—a grid of points referenced by <x, y> pairs. In most graphics technologies, the upper-left corner of the window is at (0,0) and the coordinates are positive numbers (not necessarily integers). This corner is typically the most stable, given the way window systems and interfaces handle changes to window size: dragging the lower-right corner with the mouse is probably the most common resize action across all operating systems, and that leaves the upperleft corner unaffected. Even where a system or application provides a way to resize a

window at the top or left, the contents usually stay at the resulting upper-left corner. This is illustrated in Figure 11-1.

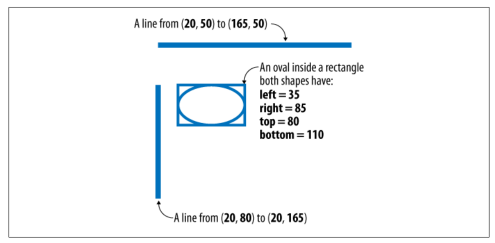


Figure 11-1. Illustration of coordinate system

Unfortunately for beginners, this coordinate system does not correspond to any of the quadrants of the traditional Cartesian system. The position of the origin relative to the quadrant corresponds to the lower-right Cartesian quadrant, in which x is positive and y negative. For designing graphical user interfaces, this can be somewhat disorienting at first. And for graphics representing quantitative data, it's really confusing, because every v coordinate must be negated. That is, the heights of titles, legends, and other parts of a diagram that are not part of the quantitative display must be *subtracted* from the y coordinates of each data point.

The basic unit of a coordinate system is the pixel. For most purposes, "pixel" means the smallest dot a device can display independently. Computer display measurements are an example: 1280×1024 means that the computer screen has 1,280 columns and 1024 rows of individually addressable points. However, all modern technologies that involve "layout"—graphics, word processing, web pages, etc.—allow measures to be specified in different units. Table 11-1 lists the measurements implemented by most systems, along with their common abbreviations. Of course, not every system supports every measure or every abbreviation.

Table 11-1. Units of measurement used in software

Name	Abbreviations	Meaning of 1 of the unit
Pixel (default)	рх	Smallest unit of the display device
Inch	i,in	Number of "dots per inch" of the display device
Millimeter	mm	Relative to inch

Name	Abbreviations	Meaning of 1 of the unit
Centimeter	CM	Relative to inch
Point ^a	p, pt	1/72 inch
Pica	P, pc	12 points, 1/6 inch
Em	em	Point size of the component's font
Ex	ex	x-height of the current font

^a "Point" and "pica" are printers' terms with a very long history and slightly different values across countries and industries. Modern computer technologies have standardized on the definition of a "point" as 1/72 inch.

A component's bounding box is the rectangle that minimally encloses the component within the coordinate system. For basic shapes it is obvious what the bounding box is, but this is not always the case for more complicated shapes. Depending on the technology, the edge of the bounding box may include one or more spaces designated as the border, padding, and margin. The "border" is a line drawn on a bounding box edge. The width of the line and its color can be specified. "Padding" usually adds extra space between what the component requires and its actual bounding box. The "margin" usually refers to extra space between a component's bounding box and a neighboring component.

Some graphics systems allow the width of each of these spaces to be specified separately for each of the four sides of the bounding box, while others allow only horizontal and vertical distances to be specified. Some systems allow a color to be specified for margins and/or padding as well as for borders. The definitions of the "height" and "width" of a component, while related to its bounding box, also differ from system to system depending on whether they include padding, borders, and/or margins.

Components

The primitive components of structured images are text items and a variety of shapes. For each, there is at least one point that specifies its location. Other points and lengths designate other aspects of its geometry. The components are defined as follows:

Text

An individual piece of text usually "wrapped" to fit within its bounding box's width Line

From one point to another

Polyline

A series of points, with a line drawn from each to the next; in effect, a composite of line segments where each one's end point is the same as the next one's start point

Rectangle

Expressed using either two opposite corners or as a corner and a width and height

Polygon

A closed shape defined by a series of points, with a line drawn from each point to the next and from the last to the first

Circle

Defined using a center point and radius length

Oval

May be defined by its rectangular bounding box or by its center point and its vertical and horizontal radii

Arc

A segment of an oval specified in one of a number of ways (e.g., specifying the oval along with a start and stop angle in degrees); whether the arc is meant to correspond to the smaller portion of the oval or the larger must also be indicated

Cubic spline

A curve defined by a start point, an end point, and one control point

Quadratic spline

A curve defined by a start point, an end point, and two control points

Composite

Several components grouped to form a new component

Composition is recursive: since composites are components, they can be included in other composites. A "top-level" component is one that is not included in a composite. Top-level components are not necessarily composites, though. For instance, a drawing may consist of a rectangle and a composite containing two lines; such a drawing would consist of three shapes and two top-level components: a rectangle and a composite.

Properties

Text and most shapes have three areas that can be controlled separately with respect to color: foreground, background, and border. Depending on the shape and the area, a "pattern"—a bitmap or image displayed repeatedly—may be used in place of a color. The color property may be called *fill* or *stroke* depending on the technology and whether or not the shape has area (a rectangle versus a line, for example).

Text objects also have fonts and other properties regular shapes do not. Some, but not all, technologies allow changes to font properties within a text object. If they do not, a composite can be used to combine two text objects with different font properties.

Other properties are specific to one or more shapes. For example, a rectangle may have a rounding angle to specify how its corners are drawn, and an arc defined as a segment of an oval will have properties such as an angle to specify what part of the ellipse it includes and a starting point along the ellipse.

Properties are generally optional when creating a component, and most properties can be modified once a component has been created (either to refine the layout while the diagram is being constructed or to add some dynamic aspects to the display).

GUI Toolkits

Window-based technologies do more than just support structured graphics. They are complete toolkits for building graphical user interfaces (GUIs).

The primary purposes of a graphical user interface are to display content—documents, images, etc.—to the user and to provide mechanisms—menus, buttons, scrollbars, input fields, etc.—for invoking application actions. The term widget is used as a general noncommittal term to refer to controls and other interface components.

Very few applications provide GUIs for structured graphics content. However, toolkits generally provide a canvas widget that supports structured drawing. Canvases can be added and displayed along with the rest of the interface and content.

GUI toolkits are not usually the best way to generate images for visualizing bioinformatics data. They often do more than is needed, and they may be difficult to learn to use. Other reasons why a GUI toolkit might not be appropriate include:

- The application may produce image files without displaying anything, often executing with no user interaction.
- The application may generate images to be displayed in a web browser, in which case the browser supplies the GUI technology.
- The application may simply need to display an image in a window, in which case you don't need buttons, menus, and all the other elaborate features of a GUI your operating system's mechanisms for managing windows are good enough.

Still, it can be useful to learn the basic details of how to display an image on the screen using a particular toolkit. Just because it provides elaborate controls and fancy features doesn't mean you have to use them.

GUI toolkits for Python

Python does not implement its own GUI toolkit. Instead, you will use one of four popular platform-independent graphical user interface libraries, along with a Python interface. This is analogous to how SQLite3 and other relational databases are used from Python. As with the sqlite3 module, the interface for one of these—the tkinter module interface to Tk—is built into Python.

The four toolkits, along with the URLs to their home pages and the corresponding Python interface libraries, are listed in Table 11-2.

Table 11-2. Cross-platform GUI toolkits and their Python interfaces

Toolkit	Home page	Python toolkit	Python interface home page
Tk	http://activestate.com/activetcl	tkinter	tkinter module documentation
wxWidgets	http://wxwidgets.org	wxPython	http://wxpython.org
Qt	http://qt.nokia.com	PyQt	http://riverbankcomputing.com/software/pyqt/intro
GTK+	http://gtk.org	PyGTK	http://pygtk.org

One or more of these toolkits may already be installed on your computer. If not, you can get installers and source code from the toolkits' home pages. As of November 2009, both tkinter and PyQt have been updated for Python 3, but wxPython and PyGTK have not. Tk itself is not part of tkinter, so if it's not on your system you'll need to go to the web page shown in the table to download an installer. Even if it is on your system, it may be an older version; if you get error messages about version incompatibilities, download and install the latest version.

A great deal of material has been published in books and on the Web about each of these cross-platform toolkits. You should be able to find tutorials, examples, forums, and many other sorts of resources that can help you get started with any of these and solve any problems you may encounter when using them later.

If you want to experiment with building applications with user interfaces, start with the tkinter documentation and the resources listed at its beginning. IDLE is implemented using tkinter. You can find its code in a subdirectory called idlelib in your Python installation's library directory. That will give you a large body of tkinter code to read.

Using a GUI toolkit

Interfaces using GUI toolkits have several high-level layers. As with structured graphics components, the details vary among toolkits. In general, there is an outermost structure called a frame that contains everything displayed—both controls and content. Around the frame is the window, managed by the operating system. Inside the frame are one or more geometry managers, or, as they are sometimes called, layout managers.

Every control and content item, including canvases, is added to a parent item, with the frame acting as the top-level parent. The layout manager automates the work of arranging their exact placement according to its policy. Policies are provided for putting each item next to the previous one either horizontally or vertically, arranging the items in a grid, and so on. Some toolkits provide a wide range of layout managers and policies, and others provide only a few. Some require layout managers to be explicitly created and added to frames, while others automatically create a layout manager for each frame. Many toolkits require a call to a function for the layout manager to do its work once its elements have all been added. Others require a call to a function for each element added.

To connect the GUI to the application, the application needs to initialize the toolkit, add the necessary elements (frames, layout managers, controls, and content), then hand control to the toolkit. As a result, GUI-based applications are largely event-driven. The toolkit waits for the user to press a key, move the cursor, or click a mouse or trackpad button. When it receives such an "event," it invokes an application action (the application establishes which action will be invoked when each control is created). The waitevent-act cycle is usually called the *event loop*, for obvious reasons, and the toolkit will include a high-level function that the application calls to start the cycle.

Using a GUI toolkit simply to display images on the screen typically requires only a few steps (and not all of these will be necessary in every toolkit):

- 1. Initialize the toolkit.
- 2. Create a frame.
- 3. Create a layout manager for the frame.
- 4. Add a canvas to the layout manager.
- 5. Create shapes and text on the canvas.
- 6. Invoke the event loop function.

Structured Graphics with tkinter

The introduction to tkinter found here is far from complete. Remember that tkinter is meant to be a full GUI toolkit, not just one for constructing graphics. The only one of its "widgets" we really need for our current purposes is its Canvas class. We will ignore controls entirely: our programs will simply put up a display and take it down when the user hits the Return key.

tkinter Fundamentals

There are several ways to organize a tkinter-based program. You can just use functions, or you can define a class. If you define a class, it can be independent of tkinter or it can inherit from Frame. Inheriting from Frame would add very little to the way we will be using tkinter, but classes are very useful in graphics programming, so we'll use independent classes.

The basic steps

The high-level steps of a minimal tkinter structured graphics program are:

- 1. root = tkinter.Tk() Initialize tkinter and get a top-level instance.
- 2. root.title(string) Give the window the title *string*.

- 3. canvas = tkinter.Canvas(root)
 - Create the canvas under the root.*
- 4. canvas.pack()

Call the Packer geometry manager to display the canvas.

- 5. Call Canvas drawing methods on the canvas.
- 6. root.close()

Close the window.

An instance of Tk represents a Tk environment. An application may have multiple instances of Tk open at the same time. As soon as a Tk instance is created, a small window appears on the screen.

There is a class for each kind of tkinter widget. The examples that follow use just Canvas. The first initialization parameter of each tkinter widget class is the existing container to which the new object should be added. This is called its "master" and must be the top-level Tk object or one of the very few other "container" components tkinter provides. The examples put the canvas directly into the top-level Tk instance.

Except for the master, all parameters in widget-creation calls are optional. These parameters specify details about the object being created. All have reasonable defaults.

Common widget options

Frequently used widget options are shown in Table 11-3. These options can be specified as keyword arguments when the widget is created, or they can be changed later with a call to the widget's configure method. Each of these options is supported by most, if not all, of the widgets.

						ptions

Name	Keyword	Value type	Units (first is default)
Background color	bg	Color value	An X11 color name or a 6-hexadecimal-digit "RGB" string (color name)
Foreground color	fg	Color value	'#RRGGBB'
Border width	bd	Distance	(px), p(t), i(n), c(m), m(m)
Width	width	Integer	Relative to width (height) of average character or widget's font
Height	height	Integer	
Font	font	Font	String or tuple; see following discussion
Anchor	anchor	Direction	'center','n','e','w','s','ne','nw','se','sw'

^{*} Either a dictionary or a series of keyword arguments may follow root to specify any of the following canvas properties: background, bd, bg, borderwidth, closeenough, confine, cursor, height, highlightbackground, highlightcolor, highlightthickness, insertbackground, and insertborderwidth (for example, tkinter.Canvas(root, bg='gray', bd=4).

The anchor parameter specifies the alignment of a widget's content within the widget. Its value indicates a location relative to the content, which will be aligned with a corresponding location in the widget. For instance, 'nw' means that the upper-left corner of the content will be aligned with the upper-left corner of the widget.

The '#RRGGBB' form of color specification works the way it does in HTML and CSS. (The three-digit form is not acceptable.) Each pair of hexadecimal digits represents an integer between 0 and 255. White would be '#FFFFFF', black '#000000', and red '#FF0000'. X11 color names come from a large set that predates the Web.† For grayscale graphics it is worth noting that the names include 'gray0' (black) through 'gray100' (white), in increments of 1.

Fonts are specified either as a family name (e.g., 'Futura') or as a tuple of the form:

```
(family, size[, variation1[, variation2[, ...]]])
```

In the tuple, *family* can be either a single family name or a sequence of family names. If a sequence, tkinter will attempt to locate each one in the operating system environment in turn until a match is found. The variations are values such as 'bold', 'italic', and 'underline'. Any number of variations may be present.

Canvas drawing methods

The Canvas class implements six drawing methods. You can use pdb to step through the drawing actions; as each item is drawn it will appear immediately in the canvas. These are not actually widgets—they are not part of tkinter's widget class hierarchy, and they are accessible only through their canvas. Within the canvas, though, they behave very much like regular widgets in most respects. The methods are:

```
create text(x, y, string[, keyword-arguments])
   Draw string at (x,y).
```

```
create line(x1, y1, x2, y2[[, . . .], keyword-arguments])
```

Draw a line from (x_1,y_1) to (x_2,y_2) and so on, for however many points there are (used for both lines and polylines).

```
create rectangle(x1, y1, x2, y2[, keyword-arguments])
```

Draw a rectangle with (x1,y1) and (x2,y2) as opposite corners.

```
create polygon(x1, y1, x2, y2, x3, y3[[, . . .], keyword-arguments])
```

Draw a polygon described by the line from (x1,y1) to (x2,y2) and then to each of the other points, finally drawing a line from the last point to the first.

```
create_oval(x1, y1, x2, y2[, keyword-arguments])
```

Create an oval inside the rectangle defined by opposite points (x1,y1) and (x2,y2).

```
create_arc(x1, y1, x2, y2[, keyword-arguments])
```

Create an arc inside the rectangle defined by opposite points (x1,y1) and (x2,y2).

[†] See http://www.tcl.tk/man/tcl8.5/TkCmd/colors.htm for a list of names and RGB values.

Table 11-4 shows some of the keyword arguments commonly used when creating and modifying objects on tkinter canvases. They are similar to those used for widgets in general, but some have different names and meanings.

Table 11-4. General canvas object properties

Name	Keyword	Value type
Inside color	fill	Color value
Border color	outline	Color value
Border width	width	Distance

Note that the width and height of a graphic object on a Canvas are not specified directly: they are derived from the x and y values in the call to the drawing method.

Each kind of drawing object also has its own specific options. Table 11-5 shows the ones we will use for text objects.

Table 11-5. Canvas text properties

Name	Keyword	Value type
Text	text	String
Font	font	Font
Anchor	anchor	Direction

Example 11-1 is a small program that demonstrates the actions to set up a tkinter window with just a canvas and an example of each kind of drawing operation. At the end, it asks the user to press the Return key, at which point it closes the window.

Example 11-1. Example tkinter program for structured graphics

```
colors = ('gray80', 'black', 'black', 'gray30',
           gray50', 'gray70', 'gray30', 'black')
import tkinter
tk = tkinter.Tk()
                                     # initialize tkinter
tk.title('demo')
                                     # give window a title
canvas = tkinter.Canvas(tk)
                                     # create the canvas
canvas.pack()
                                     # use the Packer geometry manager
canvas.create rectangle(20, 10, 120, 80, fill=colors[0])
canvas.create_line(20, 100, 120, 100, width=4, fill=colors[1])
canvas.create_line(20, 170, 45, 115, 70, 170, 95, 115,
                   120, 170, fill=colors[2])
canvas.create polygon(150, 180, 175, 115, 190, 160, 215, 125,
                       240, 180, fill=colors[3])
canvas.create oval(180, 55, 250, 115, fill=colors[4])
canvas.create arc(140, 10, 240, 80, extent=240,
                  width=4, fill=colors[5], outline=colors[6])
canvas.create text(20, 190, text='Drawing on a tkinter canvas',
                   anchor='w', font=(('Verdana, 'sans'), 14, 'italic'),
                   fill=colors[7])
```

input('Press the Return key to close the window(s)')
tk.destroy() # close the window

Figure 11-2 shows the result of running this little program.

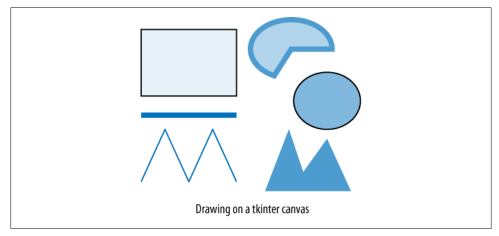


Figure 11-2. A small tkinter example

Writing the contents of a canvas to a PostScript file

I mentioned earlier that some structured graphics applications write image descriptions to files, and others write them to the screen. The Canvas class makes tkinter a bit of a hybrid, because it has the ability to write a PostScript description of its contents to a file. The PostScript file can then be sent to a printer, viewed on the screen with software that can display files in that format, or converted to another format by one of the many available image-conversion tools. A single method call suffices, specifying the filename, the width and height (in pixels) of the area to be captured, and whether the output should be in black and white (mono), gray, or color:

The background of the PostScript image will be white, ignoring the bg setting for the canvas. You may find when you view the resulting PostScript image, or a version of it converted into another format, that it is not the same size as the tkinter image was on the screen. You can scale the image proportionally in both horizontal and vertical dimensions by specifying either the pagewidth or the pageheight keyword argument, as either an int or a float (if you specify both, the value of pagewidth will be used and pageheight will be ignored). For instance, adding pagewidth=0.6*width to the preceding call would reduce the size of the PostScript image by 60% in each dimension. Note that pagewidth and pageheight represent the size of the image on the page—e.g., 800—not a percentage: the example shows how to scale an image whose width you already know.

If the purpose of your graphics program is simply to produce the PostScript file and not to display anything on the screen, then add the following after the call to canvas.postscript:

tk.destroy()

This kills the frame and all of its content. In a reasonably short program, the canvas will pack, the PostScript file will be written, and the frame and canvas will be destroyed before the window even appears on the screen. For a longer program, the frame may appear briefly in a window, though not necessarily with its canvas contents.

Examples

We're going to look now at a class that uses tkinter to draw dot plots, and another that produces histograms. Despite their different content, the steps required to draw them and many of their details are quite similar. Even where the details vary, we will be able to parameterize them. Therefore, we will first develop an abstract class that will be used as the superclass of classes implemented to draw specific kinds of visualizations.

An abstract class for tkinter graphics

Building a full-featured class that could handle a wide variety of graphical content is a far larger undertaking than would be appropriate here. Besides, such a facility would involve a number of drawing and component classes, not just a single class. The purpose of this exercise is simply to demonstrate the power of using inheritance to implement a framework (we saw an example of this with HTML.parser, in "Structured HTML parsing: html.parser" on page 298). It's also worth pointing out that one normally doesn't start by developing such a class. Rather, a particular example is developed, then another, and perhaps a third. Only as the similarities become apparent are they abstracted into a new shared superclass.

Before we look at the class itself, let's examine a short test script found at the end of its module. Example 11-2 contains the code. The class field and the two methods defined in its EmptyPlot class override those inherited from Plot. Note that since the class does not override init, the definition of init it inherits from its superclass gets called. The execute method called to display the plot is defined in the abstract class.

[‡] There is an excellent library called PyChart that produces very high-quality graphics in Encapsulated PostScript (for publishing), PDF, PNG, and SVG. It uses Python only, so it doesn't require anything else to be installed. It provides a full range of chart and support classes, making it quite easy to use to produce sophisticated graphics. Because it supports PNG and SVG, it is an excellent tool for generating graphical displays for the Web using CGI scripts. You can view examples on its creator's website at http://home.gna .org/pychart/examples/index.html. The version supplied on that website is for Python 2. You can find a version compatible with Python 3 on this book's website, along with an example application that uses PyChart to generate dot plots through a web interface.

Example 11-2. A subclass for testing an abstract class

```
if name == ' main ':
   class EmptyPlot(Plot):
       canvas background = 'gray90'
       def get plot dimensions(self):
           return 200, 150, 20, 20, 10, 10
       def draw plot(self):
           self.draw line(0, 0, 100, 100)
   plot = None
   try:
       plot = EmptyPlot()
       plot.execute()
       input(
'''You should see a light-gray plot with a line
from (0,0) to (100,100); press Return to close ''')
   finally:
       if plot:
           plot.close()
```



Something like this should go in almost every module. The code serves multiple purposes:

- Importing or executing the module will reveal syntax errors.
- Importing or executing the module can reveal basic problems with the implementation of the class, such as missing methods or modules.
- The program shows the reader the basic use of the class.
- In a graphics program, the test can display a simple diagram that shows what the class does on its own.

Detecting syntax and runtime errors is not something that you should stop doing once everything seems to work: it's a process you'll want to repeat every time you make significant changes to the class.

In this case, because the class is abstract, it is not meant to be used entirely on its own. Therefore, the test code creates a very simple subclass and uses that instead. Figure 11-3 shows what it displays. Not only does this test the code, but it gives anyone developing a subclass a guide to the abstract class's basic facilities.

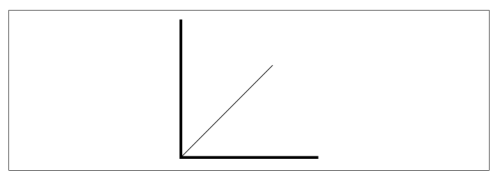


Figure 11-3. Basic example using the Plot class

The abstract class provides the following:

- Default values for class fields that control many details of the display
- An initialization method that sets off a cascade of "setup" methods that use the values defined by the class fields and the results of certain other methods
- A hierarchy of "setup" methods that defines what needs to be set up and provides reasonable defaults for some of these actions and empty methods for others
- A simplified interface to the Canvas drawing methods
- A few utilities, including one for finding fonts
- An execute method to create the widget(s), draw on the canvas, and optionally write the canvas contents out to a PostScript file

Example 11-3 shows the hierarchy of calls among the class's methods. (The actual code is available on the book's website.) What's important about this class is not its use of tkinter, but how it organizes everything for subclasses to more easily create certain kinds of plots using tkinter's Canvas drawing methods.

```
Example 11-3. The methods of the abstract Plot class
```

```
Key for subclass definitions:
   x normally would NOT implement (framework definition)
    ! MUST OVERRIDE
    * MUST EXTEND
   + MAY EXTEND
    / MAY EXTEND or OVERRIDE
    - MAY OVERRIDE (empty methods: convenience and clarity)
Utility Methods
х
   NextPlotNumber()
   closeall()
   findfont(families, size, boldflg, italicflg)
Access
   get root
   add font
```

```
get font
    __str__
Layout Framework
               tracks instance, creates Tk root, stores parameters
    init
Х
       setup
           setup fonts
           setup data
           setup parameters
           get plot dimensions
           determine layout canvas width & height, origin x,y
Drawing Framework
   execute
       create_widgets
Х
           create canvas
       draw
Х
       draw axes
Х
           draw x tics
           draw_y_tics
           draw x axis labels
           draw_y_axis_labels
!
       draw plot
   close
Drawing Functions
x draw line
x draw_line_unscaled
x draw oval
x draw rectangle
  draw text
  draw text unscaled
```

The basic steps in a program that uses this class are:

- 1. Create an instance.
- 2. Call execute and wait for the user to end the program.
- 3. Call close in the finally clause of a try statement.

The subclass is required to implement only three methods:

__init__([windowtitle[, scale[, ps_filename[, ps_scale]]]])
 which takes the following optional parameters:
 windowtitle
 A string; if false, one will be constructed
 scale

A multiplier for all <x,y> values, including the axes

ps_filename

The path to a file to which the contents of the canvas in PostScript can be written

ps scale

The scale to use for the PostScript version of the drawing

get plot dimensions()

which must return a tuple of the following form containing the details of the plot within the canvas:

(width, height, leftpad, rightpad, toppad, bottompad)

width and height are the dimensions of the area of the plot itself—i.e., the area defined by the extent of the x and y axes. The other values indicate how much space to leave between the corresponding edge of the canvas and the plot. These values, along with the values of certain class fields, are used in determine layout to set a number of values used in organizing the display.

• draw plot()

which creates the objects that go in the Canvas and specifies their properties.

The subclass is free to implement setup data and setup parameters however it wishes. It is intended that one or both of these methods, or perhaps just the subclass's init method, will determine and store the values that will be used in draw plot. Note that draw plot is not responsible for drawing axis lines, tics, or labels. The drawing functions are relative to an origin at the *lower left*, with positive values above the axis. This inverts the usual direction of y coordinates in tkinter to make plotting more convenient.

Table 11-6 shows the class fields that control details of the display. Any of them may be overridden in a subclass. Figure 11-4 illustrates the basic layout implemented by the drawing framework. The four margins and the plot height and width are values returned by a subclass's get plot dimensions method. The two axis widths are the values of class fields. The diagonal drawn demonstrates that y values increase upward. A line like this, along with the circle and text at its end, is drawn using Plot's drawing methods. Positions are expressed relative to the origin of the plot area, not the canvas.

Table 11-6. Class fields used by Plot class to control display

Name	Value	Use
PlotName	'Plot'	$\label{lem:combined} Combined with the current instance count to construct a default window name$
canvas_pad_x,	0	Horizontal and vertical padding of the canvas widget itself
canvas_pad_y		
canvas_border	0	Border width of the canvas
canvas_background	'white'	Background color of the canvas
x_axis_width	2	Width of the axis lines; o means "no line"
y_axis_ width		
x_tic_length	12	Length of "tic" lines on axes if axes are drawn

Name	Value	Use
y_tic_length		
<pre>x_axis_font_size y_axis_font_size</pre>	12	Size of font for labels drawn on axis, if any
serif_faces	Lists of likely font names	Convenient defaults for subclasses to use as desired
sans_faces		
mono_faces		

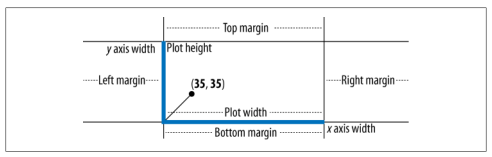


Figure 11-4. Drawing framework layout

Dot plots

Dot plots are a simple technique for visually comparing two base or amino acid sequences of similar size to highlight slight differences. One sequence is arranged along the horizontal axis and the other along the vertical axis. Then, the base or amino acid at each sequence on one axis is compared to every one on the other. In the simplest implementation, the plot contains a dot wherever there is an exact match. Figure 11-5 illustrates the process with two short sequences.

Due to random matches, however (more so in base sequences than in amino acid sequences), this implementation generates far too many dots to be useful for many purposes. Random matches can be reduced by using two parameters: a window w and a cutoff c. At each point, that point plus the next w-1 points are compared, and the total number of matches is counted. A match is registered for that point only if the count is at least c. A graphical display of the results could use multiple levels of gray to show different scores above c. For some applications, amino acid sequence matching can be refined further by scoring each comparison according to a substitution matrix rather than just testing equality.

[§] A good review article that covers the history, uses, and algorithms of two-sequence dot plots in bioinformatics can be found at http://www.code10.info/index.php?view=article&id=64. It includes explanations of how the various kinds of differences between a pair of sequences appear in dot plots.

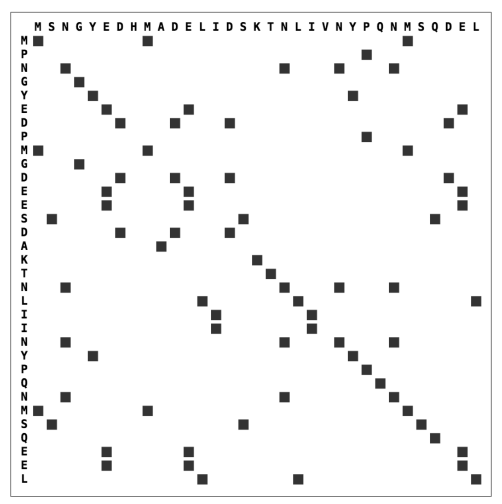


Figure 11-5. Comparison of two sequences for a dot plot

Figure 11-6 illustrates the effects of these parameters for a short amino acid sequence compared against itself. On the left, w and t are both 1. On the right, w=3 and t=2. These are more typical values for amino acid sequences (typical values for base sequences would be w=11 and c=7). The actual values to use for these parameters will depend on the purpose of the comparison. An application could easily provide controls that allow the user to see what the dot plot looks like for different values of w and t.

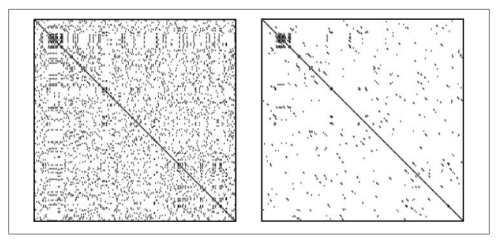


Figure 11-6. A dot plot with different windows and cutoffs

A perfect match between two sequences will show a solid diagonal line. Two sequences that match exactly except for their initial parts will also show a solid diagonal line, but unless the nonmatching parts are the same length the diagonal will not be along the x=y diagonal of the graph. Dots scattered elsewhere are not meaningful unless clustered in certain ways.

Dot plots can visually highlight evolutionary modifications both between two similar sequences and within a sequence. The effects of the possible changes are listed in Table 11-7. Some applications display dot plots with the diagonal going from upper-left to lower-right, and others from lower-right to lower-left. For this reason, the table just says "toward the right" or "toward the left." The direction is to be interpreted as relative to the x=y diagonal.

Table 11-7. Appearance of mismatches in dot plots

Change in y-axis sequence	Visual effect
Mutation	Small gaps in the line
Deletion	Line offset to the right of the diagonal
Insertion	Line offset to the left of the diagonal
Duplication	Line offset to the left of the diagonal, with a parallel overlap corresponding to the duplication
Inversion	Large gap with a faint, broken line of similar length perpendicular to the gap and centered on its center
Self-duplication	Two line segments reflected around the diagonal when the sequence is compared to itself

Two other patterns have special significance. One is an artifact of sequencing technology: a stretch of Ns in one sequence where the other sequence has real bases will result in a horizontal or vertical swatch without dots.

The other pattern does have biological significance, but not necessarily an evolutionary one: an area of low complexity manifests itself as dense, perhaps even solid, squares around the diagonal. In base sequences, these squares reveal areas with many repeats of a very short subsequence. An amino acid sequence would usually not have multiple repeats of exactly the same short subsequence, but a stretch containing only a few of the possible amino acids would have a similar effect (e.g., EKLKEKEKQKEKERQREKEK). These squares can even reveal regions of self-similarity when a sequence is compared with itself.

Figures 11-7, 11-8, and 11-9 illustrate the way the changes shown in Table 11-7 appear in dot plots. These illustrations were constructed manually by making changes in a copy of the x-axis sequence, an amino acid sequence of length 292. (Dot plots are often used for much larger sequences, of course, such as cDNA sequences.)

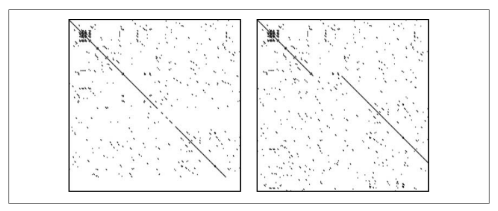


Figure 11-7. Mutations and deletions in a sequence dot plot

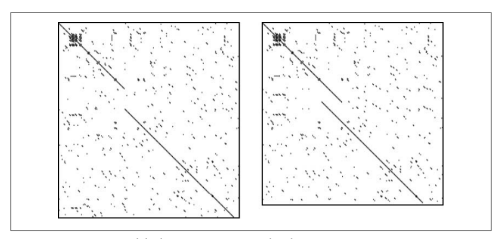


Figure 11-8. Insertion and duplication in a sequence dot plot

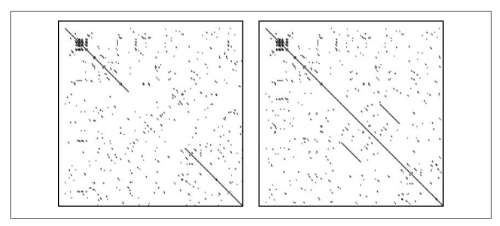


Figure 11-9. Inversion and self-duplication in a sequence dot plot

Next, we'll look at a complete definition of a class for generating dot plots. It is based on the abstract Plot class described earlier. Where methods are defined that override or extend the inherited versions, the method name is highlighted. In addition, if the method extends the inherited method, the call to super is highlighted.

In anticipation of its use in real applications, our class provides a class method that returns an optparser facility. One of its __init__ method's required arguments is an OptionParser that has already parsed the command line. Quite a few parameters are set through the parser. Rather than passing them separately when the DotPlot is created, the option object packages them. For convenience, methods throughout the class are defined to access fields of the OptionParser pretty much as if they had been fields of the DotPlot object.

The module includes test code just the way the abstract class's module did, as shown in Example 11-4. It generates a random sequence 201 amino acids long and plots it against itself. Figure 11-10 shows the plot displayed.

Example 11-4. Test code for the DotPlot module

```
except:
                                                      # 110
    input("Press the Return key to close the window(s)")
    plot.close()
```

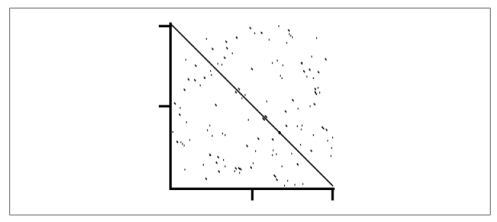


Figure 11-10. The plot drawn by the DotPlot test code

Example 11-5 is the first part of the class definition. It overrides a couple of class fields and defines similar parameters for its own use. The init method it shows is unusual in that it calls its superclass's <u>_init</u> last rather than first. This is because the abstract class's init method sets off a cascade of initialization and setup methods that will use some of the values stored by this class's init method.

```
Example 11-5. Generating a dot plot using tkinter
```

```
class DotPlot(Plot):
# Overridden class fields
    x tic width = y tic width = 3
    sans faces = ('Helvetica Narrow', 'Futura',
                   'Helvetica', 'Arial', 'sans-serif')
    PlotName = 'Dot Plot'
# Fields specific to this class
    title font size = 12
    top title margin = round(title font size / 2)
    name height = 22
    tic_width = 3
    axis width = 3
    # Defaults for showing the plot without axes;
    # if options.axes is true, they will be reset in setup_parameters
    left_title_margin = 10
    xaxis width = yaxis width = 0
    xlabel height = ylabel width = 0
    def init (self, seq1, seq2, seqname1 = '', seqname2='',
                  window=1, threshold=1, with_axes=False, dot_size=1,
                  # super parameters:
```

```
window_title=None,
    scale=1.0, ps_filename=None, ps_scale = 1.0):
self.seq1 = seq1
self.seq2 = seq2
self.seqname1 = seqname1
self.seqname2 = seqname2
self.window = window
self.threshold = threshold
self.with_axes = with_axes
self.dot_size = dot_size
self.window_title = window_title
super(). init_(options.name, scale, ps_filename, ps_scale)
```

Example 11-6 contains configuration methods that extend or replace definitions inherited from Plot. These mostly determine details that are specific to the kind of plot the class implements.

Example 11-6. Plot setup methods overridden in DotPlot

```
def setup fonts(self):
    super().setup fonts()
    self.add font('title', self.findfont(self.sans faces,
                                         self.title font size))
def setup data(self):
    self.points = self.compute points()
    self.max x = max(self.points, key=lambda pt: pt[0])[0]
    self.max y = max(self.points, key=lambda pt: pt[1])[1]
def setup parameters(self):
    super().setup parameters()
    if not self.with axes:
        self.x axis width = self.y axis width = 0
    else:
        self.x axis width = self.axis width
        self.y axis width = self.axis width
        self.x label height = self.x tic length + 5
        self.y label width = self.y tic length + 5
        self.left title margin = \
            self.left title margin + self.y label width
def get plot dimensions(self):
                                              # required
    return (round((self.max x + self.window) * self.scale),
             round((self.max y + self.window) * self.scale) +
             self.y_tic_width,
             self.y_label_width,
             self.ipad right,
             self.name height *
             (bool(self.seqname1) + bool(self.seqname2)),
             self.x label height)
```

Example 11-7 shows the code that determines the points to be plotted. The setup_data method from Example 11-6 calls compute_points and assigns the points field to its result.

```
def compute points(self):
    pts = []
    for y in range(1 + len(self.seq2) - self.window):
        for x in range(1 + len(self.seq1) - self.window):
            if self.test point(self.seq1, x, self.seq2, y):
                pts.append((x, y))
    return pts
def test point(self, seq1, x, seq2, y):
    cnt = 0
    for n in range(self.window):
        if seq1[x+n] == seq2[y+n]:
            cnt += 1
    return cnt >= self.threshold
```

Example 11-8 implements the class's drawing operations. If sequence names were provided in the call to DotPlot, their names are drawn above the plot. If the with axes argument is false, the setup parameters method from Example 11-6 sets the width of both axes to 0, and Plot does not draw the axes. If it's true, setup parameters sets the axis widths along with some other parameters. In that case, the superclass draws the axis lines and calls draw x axis labels and draw y axis labels. As defined, these functions just draw tic marks every 100 points along each axis. Finally, a circle (oval) of diameter dot size pixels is drawn for each of the computed points. Few graphical user interface facilities actually provide a point-drawing operation; instead, programs simply draw very small circles.

Example 11-8. DotPlot's drawing methods

```
def draw(self):
    super().draw()
    self.draw_titles()
def draw titles(self):
    if self.segname1 and self.segname2:
        self.draw title('x = ' + self.seqname1,
                        self.top title margin)
        self.draw title('y = ' + self.seqname2,
                        self.top title margin +
                        self.title font size + 6)
    elif self.seqname1 or self.seqname2:
        self.draw title(self.seqname1 or self.seqname2,
                        self.top title margin)
def draw title(self, title, ypos):
    self.draw text unscaled(self.left_title_margin,
                            ypos, title, 'title', 'nw')
def draw x axis labels(self):
    for x in range(100, self.plot width+1, 100):
        self.draw_line(x, -self.x_axis_width,
                       x, -(self.x axis width + self.x tic length),
```

```
self.x tic width)
def draw y axis labels(self):
    adjust = round((self.y tic width - 1)/2)
    for y in range(100, self.plot height+1, 100):
        self.draw line(-self.y axis width,
                       y + adjust,
                       -(self.y axis width + self.y tic length),
                       y + adjust,
                       self.y tic width)
def draw plot(self):
                                                 # required
   for pt in self.points:
        self.draw oval(
            pt[0],
            self.plot_height - self.window - pt[1],
            pt[0] + self.dot size - 1,
            self.plot height - self.window - pt[1] - self.dot size - 1)
```

Histograms

An interesting use of histograms is to plot the distribution of codon usage among the known coding sequences for an organism. The program shown in this section color-codes the bars according to the GC content of the corresponding codon. The GC content for all of the cDNA sequences from which the data was taken is written at the upper left of the graph. (Note that this is not the same as the GC content for the organism's DNA as a whole, because it is based on coding sequences.) The code to extract the data from a file is not shown here, but it is quite similar to our many FASTA examples.

Histograms for two different bacteria are shown in Figures 11-11 and 11-12. Looking at the two figures, it's easy to see that the distributions of codons used by these two species are substantially different.#

Example 11-9 shows the class fields of another subclass that uses the abstract Plot class shown earlier. Method definitions are shown in subsequent examples. In those examples, the names of methods that override or extend methods inherited from Plot are highlighted, as are calls to super.

[∥] The program's data source is a file of bacterial species codon distributions downloaded from <code>ftp://ftp.kazusa.or.jp/pub/codon/current/gbbct.codon</code>. Interesting interactive tools for getting codon counts in several formats for species matching a regular expression can be found at <code>http://www.kazusa.or.jp/codon</code>.

[#]The downloaded data at the previously mentioned site was computed from the GenBank flat file release of June 15, 2007, using the cDNAs available for each species. The datafile simply contains alternating lines of species information and 64 integers representing counts per thousand of each codon. The program doesn't do anything to compute those values; it just graphs them.

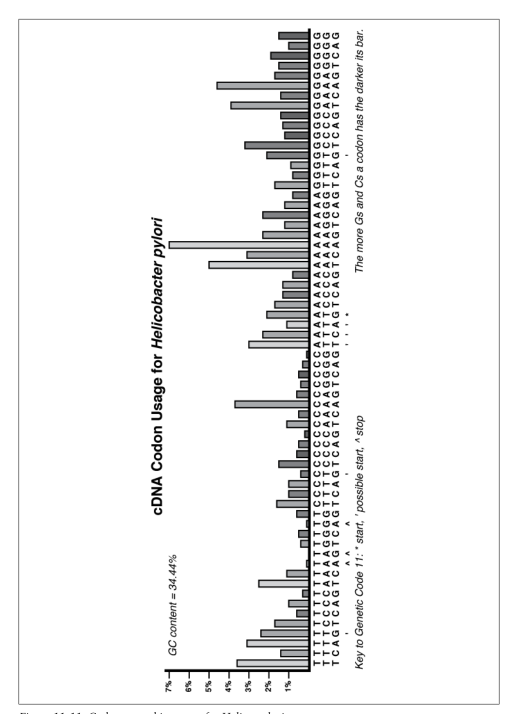


Figure 11-11. Codon usage histogram for Helico pylori

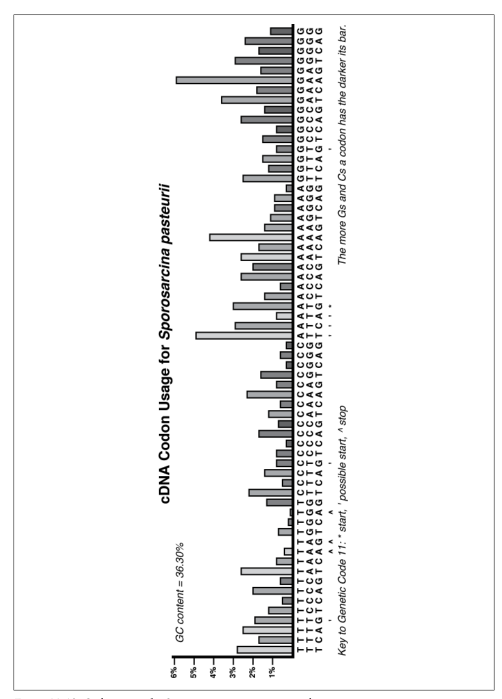


Figure 11-12. Codon usage for Sporosarcina pasteurii using tkinter

Example 11-9. Codon distribution histograms using tkinter

class CodonUsageHistogram(Plot): # Fields that override inherited values x tic length = y tic length = 16 # Fields specific to this class barwidth = 10 # width of histogram bars barspacing = 5 # spacing of histogram bars barscale = 3 # vertical pixels per data point codon font size = 14 marker font size = 14 title_font_size = 22 key font size = 16 codon faces = ('Liberation Mono Regular', 'Lucida Sans Typerwriter', 'DejaVu Sans Mono', 'Bitstream Sans Mono', 'Courier') gc colors = ('gray80', 'gray60', 'gray40', 'gray25') default bar color = 'gray94' plot top margin = 20 ylabel width = 60 x label height = codon font size*5 canvas_background = 'white' title pad y = 6key pad y = 12codons = [base1 + base2 + base3 for base1 in 'TCAG' for base2 in 'TCAG' for base3 in 'TCAG'] codon labels = codons[:] # a copy to modify startchar = '*' altstartchar = "'" stopchar = '^' gc colors = ('gray80', 'gray60', 'gray40', 'gray25') # add start and stop indicators # stop codons: for codon in ('TAA', 'TAG', 'TGA'): codon labels[codon labels.index(codon)] += stopchar # standard start codon: codon labels[codon labels.index('ATG')] = 'ATG' + startchar # alternate start codons in code 11 for codon in ('TTG', 'CTG', 'GTG', 'ATT', 'ATA', 'ATC'): codon labels[codon labels.index(codon)] += altstartchar



Notice the for statements in the class field section. Remember that when a class statement is executed, each of the statements it contains is executed. Although normally classes contain just assignment and def statements, they can contain loops, iterations, conditionals, and so on. These for statements and associated assignments set up a list of codon labels that incorporate start and stop codons as well as the alternate start codons for the bacterial genetic code.* (The class could easily be generalized to allow specifying a code as an argument when creating an instance; what's shown here assumes bacterial sequences.)

Example 11-10 shows the definition of __init__. The name parameter is the name of the organism, for the purpose of displaying a title. The data is simply a dictionary or list of pairs that gives the count per thousand of the corresponding codon. The other methods in the example are part of the setup area of the framework.

Example 11-10. Setup methods for the tkinter codon distribution plot

```
def init (self, name, data,
             # super paramaters:
             windowtitle=None, scale=1.0,
             ps filename=None, ps scale = 1.0):
    self.data = dict(data)
    self.name = name
    super(). init (name, scale, filename, ps scale)
def setup fonts(self):
    super().setup fonts()
    self.add font('x', self.findfont(self.codon faces,
                                     self.codon font size,
                                     True))
    self.add_font('title', self.findfont(self.sans_faces,
                                         self.title font size,
                                         True))
    self.add font('key', self.findfont(self.sans faces,
                                          self.key font size,
                                          False, True))
    self.add_font('marker', self.get_font('x').copy())
    self.get_font('marker').configure(size=self.marker_font_size)
    self.add font('species', self.get font('title').copy())
    self.get font('species').configure(slant='italic')
def setup data(self):
    self.maxval = max(self.data.values())
    self.sumval = sum(self.data.values())
    self.maxpercent = round((100 * self.maxval) / self.sumval)
def get plot dimensions(self):
                                                                    # required
    return (self.barspacing + round(64 * self.scale *
```

^{*} All the alternate start codons documented for the bacterial genetic code at http://www.ncbi.nlm.nih.gov/Taxonomy/Utils/wprintgc.cgi#SG11 are included, even those found very rarely.

```
(self.barwidth + self.barspacing)),
            round(self.scale * self.barscale * (2 + self.maxval)),
            self.ylabel width + 20,
            self.get title height(),
            self.xlabel height + self.get key height()
def get title height(self):
    return 2 * self.title font size
def get key height(self):
    return 2 * self.key font size
```

Example 11-11 shows the definitions of the methods that override the empty superclass methods for drawing the tics and labels of the y-axis and the elaborate codon labels along the x-axis.

Example 11-11. Axis methods for the tkinter codon distribution plot

```
def draw y tics(self):
    for n in range(1, self.maxpercent + 1):
        vertpos = 10 * n * self.barscale
        # * 10 because barheights are per thousand
        self.draw line(0, vertpos, -self.y tic length, vertpos)
def draw y axis labels(self):
    for n in range(1, self.maxpercent + 1):
        self.draw text(-self.tic length-4,
                        10 * n * self.barscale,
                        str(n) + '%'
                         'y',
                         anchor='e')
def draw x axis labels(self):
    curx = self.barspacing + 5
    for n, codon in enumerate(self.codon labels):
        self.draw codon label(curx, codon),
        curx += self.barwidth + self.barspacing
def draw codon_label(self, x, codon):
    for n, char in enumerate(codon):
        self.draw text(x,
                        # 6 * is additional separation for markers
                         -((self.codon font size * (n+1)) +
                           (6 * (n > f2))),
                        'x' if n < 3 else 'marker',
                       anchor='center')
```

The methods in Example 11-12 draw the title above the plot, the key below, and an indication of the total GC content of the coding sequences at the upper left of the plot area. The computation of the total GC content is done by the gccontent method shown in the subsequent example.

```
def draw(self):
    super().draw()
    self.draw title()
    self.draw gc content()
    self.draw key()
def draw title(self):
    self.draw text(round(self.plot width / 2), self.plot height,
                   'cDNA Codon Usage for ', 'title', anchor='se')
    self.draw text(round(self.plot width / 2), self.plot height,
                   self.name, 'species', anchor='sw')
def draw gc content(self):
    self.draw_text(20, self.plot_height-20,
                   'GC content = {:.2f}%'.format(self.gccontent()),
def draw key(self):
    self.draw text(
        0, -self.xlabel height,
        'Key to Genetic Code 11: ' +
        '{} start, {} possible start, {} stop'.
        format(self.startchar, self.altstartchar, self.stopchar),
        'key', anchor='nw')
    self.draw text(
        self.plot width, -self.xlabel height,
        'The more Gs and Cs a codon has the darker its bar.',
        'key', anchor='ne')
```

The methods in Example 11-13 draw the usage bars. Each bar is drawn in one of four shades of gray, according to the number of Gs and Cs in the codon. The two class methods are class methods simply because nothing they do is specific to the instance. The gccontent method calls codon gc count as part of its computation of the total GC content for the sequences. That is not a class method, because its computation depends on the instance's data field.

Example 11-13. Methods for drawing the distribution bars

```
@classmethod
def codon gc count(self, codon):
    return codon.count('G') + codon.count('C')
@classmethod
def gccolor(self, codon):
    return self.gc colors[self.codon gc count(codon)]
def gccontent(self):
    # 3 because there are 3 bases in each codon
    # 10 because the counts are per thousand
    return (3 * 10 * sum((self.codon gc count(codon) * count
                       for codon, count in self.data.items())) /
```

```
self.sumval
def draw plot(self):
                                            # reauired
    curx = self.barspacing
    for codon in self.codons:
        n = self.data[codon]
        self.draw rectangle(curx, 0, self.barwidth, n * self.barscale,
                            self.gccolor(codon))
        curx += self.barwidth + self.barspacing
```



The code in this chapter has a very different "shape" than other Python code shown in this book. While the overall organization of graphics programs can make good use of object-oriented techniques, in the end drawing comes down to a series of commands to draw specific things with specific properties in specific places. This results in many procedural-style function calls with many arguments.

You might also have noticed some funny arithmetic going on: adjustment factors, adding 1 or 2, calls to round, and so on. These are not typically manipulations you can anticipate, and they won't necessarily be the same if you implement the same program with a different toolkit. Rather, they come from seeing the results of straightforward calculations and then adjusting them. It is always best to at least name such values in a way that indicates their purpose, such as adjustment or border-displacement. Comments are more useful in this kind of programming than they are in more straightforward code.

Structured Graphics with SVG

SVG—Scalable Vector Graphics—defines a set of XML markup tags and attributes for describing shapes and text to be displayed in a web browser.† The name of the technology emphasizes several important characteristics:

Vector- versus pixel-based graphics

Most image formats represent pixels. They contain no information about what the pixels represent, so transformations such as zooming in and out can operate only on pixels. In contrast, a vector-based system describes what is in the image. The description is used to generate pixels when the image is displayed. (This is analogous to the difference between representing fonts as bitmaps versus representing them as unit-independent curves.)

[†] See http://www.w3.org/TR/SVG11 for the complete specification. Much of it is very formal and sophisticated, but the sections on shapes (http://www.w3.org/TR/SVG11/shapes.html) and text (http://www.w3.org/TR/ SVG11/text.html) are very useful references for even basic use of SVG. Both have excellent examples, including the resulting images, and would constitute an excellent supplement to the brief descriptions in this chapter.

Scalable technology

The term "scalable" in SVG means two things. The first one, which we won't talk about here although it's important in other contexts, is that the technology itself is scalable: one SVG file can refer to many others to build highly complex images and can be used by many different kinds of applications.

Scalable graphics

What interests us here is that a vector-based image is in itself "dimensionless." The actual image is generated for a specified resolution. This means the images can be zoomed in and out and used at different sizes on different web pages. A scalable representation supports the different requirements of different output media—i.e., the widely varying resolutions of different makes and models of displays and printers.

SVG is quite powerful: it supports scaling, resizing, interaction, and animation. An SVG file can include Cascading Style Sheets (CSS) and scripts (JavaScript). SVG content can be delivered to a browser in several forms, including:

- In a file containing just SVG
- By URL reference to a separate SVG file in an HTML or XHTML file <object> or tag
- Directly inside an XHTML file

For simple situations, a self-contained SVG file is fine. If the image is only a part of a web page to be displayed, using an object or img tag is appropriate. More sophisticated client-side programming techniques are best supported by embedding the SVG content inside an XHTML file. We will use a plain SVG file here.

SVG is generally well supported by modern browsers. In the same way that a browser interprets HTML to display a web page and other graphics formats—such as PNG, JPEG, or GIF—to display an image, it interprets the contents of an SVG file and generates the corresponding image to display. A browser's zoom commands should zoom in and out on the image, and if the entire image doesn't fit in the window the browser should show scroll bars. (Actual behavior varies among browsers.)

SVG File Contents

The SVG File template shows the general outline for an SVG file.



The "standalone" on the first line means the file refers to outside documents, which it does in the first few lines. It does not mean that this is an independent SVG file as opposed to one referenced by or embedded in another web page.

The width and height parameters can be specified as pixels (e.g., 400) or percentages followed by a percent sign (e.g., 100%). There are a number of other possibilities as well, but those will serve our purposes.

The template shows where stylesheet information and the actual SVG tags belong. We'll see examples of these a little further on in the chapter.

```
TEMPLATE
                                         SVG File
The outline of an SVG file is:
     <?xml version="1.0" standalone="no"?>
     <!DOCTYPE svg PUBLIC "-//W3C//DTD SVG 1.1//EN"
     "http://www.w3.org/Graphics/SVG/1.1/DTD/svg11.dtd">
     <svg xmlns='http://www.w3.org/2000/svg' version='1.1'</pre>
          width='width' height='height'>
        <style type='text/css'><![CDATA[</pre>
           ... CSS style rules here, just as in HTML ...
         </style>
       </defs>
       ... SVG tags ...
     </svg>
```

SVG tags

In its complete form, SVG is a sophisticated markup language. For many purposes, though, all that's needed is a simple repertoire of tags and a few of their more common attributes. Tables 11-8 (shapes) and 11-9 (text) describe those. In addition, all of the tags support the style attribute, which is described next. In the following, note that y coordinates start with 0 at the top of the diagram and increase downward. The default for all unspecified coordinates is 0.

Table 11-8. SVG shape tags

Tag	Attributes	Meaning
rect	x	Smaller of the two x-axis coordinates
	у	Smaller of the two y-axis coordinates
	width	A positive value
	height	A positive value
	rx	x-axis radius of rounded corners
	ry	y-axis radius of rounded corners
circle	сх	x-axis coordinate of center

Tag	Attributes	Meaning
	су	y-axis coordinate of center
	r	Radius
ellipse	СХ	x-axis coordinate of center
	су	y-axis coordinate of center
	rx	x-axis radius
	ry	x-axis radius
line	x1	x-axis coordinate of starting point
	у1	y-axis coordinate of starting point
	x2	x-axis coordinate of ending point
	y2	y-axis coordinate of ending point
polyline	points	Quoted list of whitespace-separated points, each a comma-
(open)		separated x,y pair
polygon	points	Quoted list of whitespace-separated points, each a comma-
(closed)		separated x,y pair

Table 11-9. SVG text tag

Tag	Attributes	Meaning
text	x	x-axis coordinate of start of text
	у	y-axis coordinate of start of text
	width	Width of text box (optional)
	height	Height of text box (optional)

SVG styles

SVG tags can be styled using CSS, just the way HTML tags can be. Two attributes, shown in Table 11-10, can be used to specify a tag's style. In addition, styles can be defined within the style tag in the defs section at the beginning of an SVG document.‡

Table 11-10. SVG style attributes

Attribute	Effect	Value types
style	Specifies the style for the tag	Semicolon-separated name: value pairs
class	Applies the style for the class as defined in the style section	A name
id	Applies the style for the elements with this id as defined in the style section	A name

[‡] Full details regarding the use of CSS with SVG can be found at http://www.w3.org/TR/SVG11/styling.html #StylingWithCSS.

The basic style properties are described in Table 11-11. Table 11-12 shows the properties used to specify fonts.§

Table 11-11. Major SVG style properties

Property	Effect	Value type
fill	Inside color	Color (name ^a , #rgb, etc.)
stroke	Border color	Color (name, #rgb, etc.)
stroke-width	Width of line	Number
transform	Modifies element (explained in following discussion)	Space- or comma-separated transforms in function-like notation

^a See http://www.w3.org/TR/SVG/types.html#ColorKeywords for a list of acceptable color names and their RGB equivalents.

Table 11-12. SVG font properties

Property	Effect	Value type
font-family	Typeface	Comma-separated list of font names to try, in the order in which they appear. Names containing whitespace must be quoted. serif, sans-serif, and monospace refer to the corresponding browser settings.
font-size	Size	Number representing points.
font-style	Italicization	Normal, italic, or oblique.
font-weight	Boldness	Normal, bold, bolder, or lighter.

The transformation sublanguage provides a powerful range of capabilities for modifying shapes and text. These include the following:

translate(tx [ty])

Add tx to the element's x position and ty (0 if omitted) to its y position.

scale(sx [sy])

Scale the element by a factor of sx horizontally and sy (sx if omitted) vertically.

rotate(angle [cx cy])

Rotate by *angle* around the point $\langle cx, cy \rangle$ (<0, 0> if omitted, which usually has very surprising effects, if you even see the object after it has been rotated).

In addition to the tags for shapes, paths, and text, SVG provides a grouping tag, g. This is similar to the div tag in HTML. It provides a way to apply styles and transformations to all the tags between the opening <g> and the closing </g>. The style for the group can be specified in all the usual ways: as the value of a style attribute, with a class attribute that corresponds to a style class, or with an id that corresponds to a style defined for that ID.

§ See http://www.w3.org/TR/2008/REC-CSS2-20080411/fonts.html#font-specification for full details.

http://www.w3.org/TR/SVG11/coords.html#TransformAttribute contains the full documentation.

Examples

Next, we'll look at two examples of programs that produce structured graphics using SVG. The first repeats the tkinter histogram example without the benefit of a graphics library. Because SVG files are ultimately just XML text, it is straightforward to generate them with ordinary Python code. For the same reason, though, SVG is less useful for graphics composed primarily of large numbers of individual points, such as the dot plots we saw earlier. Each point would require a separate SVG tag, and there could be tens of thousands of points in a real dot plot.

Histograms of codon use

Example 11-14 shows the outline of a program for producing the same sort of histograms as in the earlier tkinter examples. Example 11-15 shows the resulting SVG text. The code isn't shown here, since it is somewhat long and doesn't contain anything we haven't already seen; it is, however, available on the book's website.

Example 11-14. The outline of the codon usage histogram program

```
chart usage
   draw chart
        draw heading
        draw key
            draw text
        draw title
            draw text
        draw axes
            draw line
            draw text
        draw codon label
        draw bars
            draw bar
        draw closing
```

Figures 11-13 and 11-14 correspond to Figures 11-12 and 11-11. There are slight differences, largely because of the way the x-axis was labeled with codons. In the tkinter implementation, separate calls to Canvas.create text were used to stack characters vertically. Using SVG would have been fairly awkward, but SVG's transforms made it easy to rotate the text.



Although browser support for SVG has come a long way, browsers don't generally implement every possible feature and sometimes don't get the features they do implement quite right. It would be worth viewing the SVG files for this example in more than one browser.

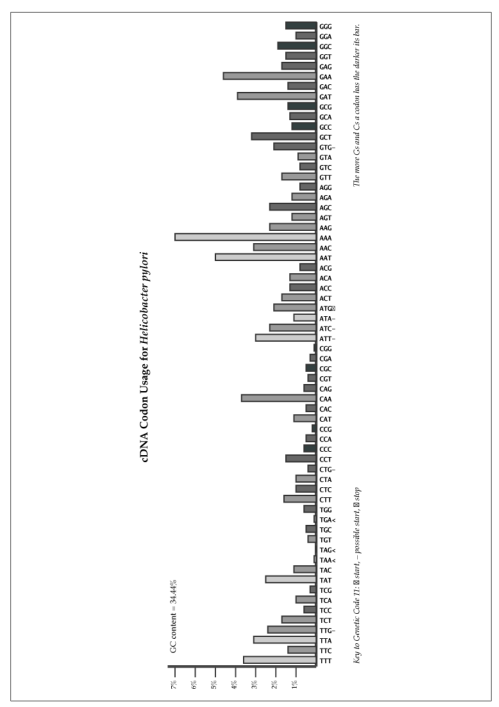


Figure 11-13. Chart of Helicobacter pylori codon usage

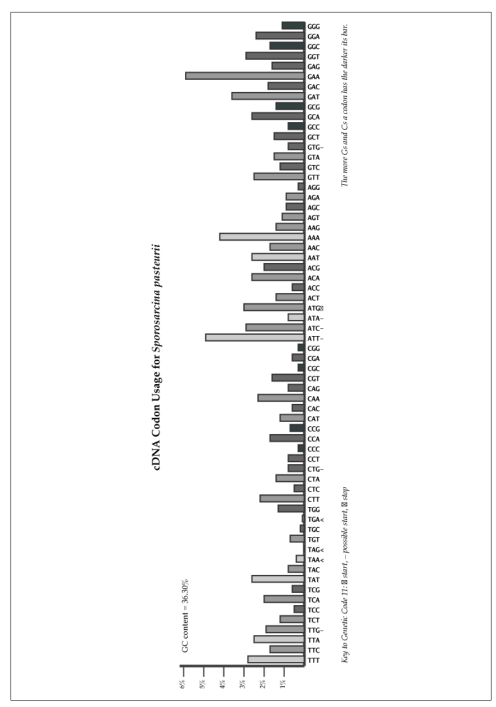


Figure 11-14. Chart of Sporosarcina pasteurii codon usage

The opening part of the SVG file generated by this example, containing the required invocations, is shown in Example 11-15. Next is a set of CSS style declarations. The details are omitted here, but the beginning and end are shown to indicate how to include CSS styles in an SVG document. (Example 11-15 includes a small section with actual CSS declarations.) The rest of the SVG file relies on these CSS declarations to describe things like color and font properties. A CSS declaration can specify properties in a variety of ways, including for a type of tag (e.g., text) or for all tags with a certain value of the class attribute. Finally, a few of the graphical element tags from each part of the file are included, with ellipses indicating more of the same.

Example 11-15. SVG for dispaying a codon usage bar graph

```
<?xml version="1.0" standalone="no"?>
<!DOCTYPE svg PUBLIC "-//W3C//DTD SVG 1.1//EN"
"http://www.w3.org/Graphics/SVG/1.1/DTD/svg11.dtd">
<svg xmlns='http://www.w3.org/2000/svg' version='1.1'</pre>
     width='984' height='379'
    <style type='text/css'><![CDATA[
        ...CSS styles go here...
     ]]>
   </style>
 </defs>
 <!-- Key -->
   <text x='100' y='339' class='key' >
      Key to Genetic Code 11:
      ✱ start, – possible start, ∧ stop
   </text>
   <text class='keyright' x='944' y='310' >
      The more Gs and Cs a codon has the darker its bar. </text>
   <text x='100' y='72' >GC content = 36.30% </text>
 <!-- Title -->
   <text class='title' x='492' y='30' >
      cDNA Codon Usage for
      <tspan font-style='italic'
             text-anchor='middle' dominant-baseline='middle' >
        Sporosarcina pasteurii
      </tspan>
    </text>
 <!-- Axes -->
   <line class='axis' x1='80' y1='246' x2='944' y2='246' />
    cline class='axis' x1='80' y1='60' x2='80' y2='246' />
  <g class='tic'>
    <line x1='64' y1='216' x2='80' y2='216' />
   <text x='60' y='217' > 1% </text>
line x1='64' y1='186' x2='80' y2='186' />
```

```
<text x='60' y='187' > 2% </text>
</g>
<!-- Codon Labels -->
<g class='codon'>
  <text x='191.5' y='251' transform='rotate(90,191.5,251)' >TAT</text>
  <text x='205.0' y='251' transform='rotate(90,205.0,251)' >TAC</text>
  <text x='218.5' y='251' transform='rotate(90,218.5,251)' >TAA&lt;</text>
  <text x='232.0' y='251' transform='rotate(90,232.0,251)' >TAG&lt;</text>
</g>
<!-- Bars -->
  <rect class='gc0' x='83.5' y='162' width='10' height='84' />
<rect class='gc1' x='97.0' y='195' width='10' height='51' />
<rect class='gc0' x='110.5' y='171' width='10' height='75' />

  <rect class='gc1' x='124.0' y='189' width='10' height='57' />
</svg>
```

Sequencing trace file curves using SVG

Many curves in bioinformatics data presentations are highly irregular. This makes them difficult to draw using typical drawing facilities, such as those tkinter provides, or a description language such as SVG. A simplifying trick that often works for these kinds of curves is to draw a line from each point in the data to the next. The lines will be so small that the overall image will appear to be a curve.

A classic example is the display of trace data obtained from sequencing machines. One approach using SVG is simply to insert a line tag from every <x, y> position to the corresponding <x+1, y'> position. It turns out that the lines drawn are so small that the end result looks like a curve.

A file containing enough line tags for a sequencing trace will be quite large. Worse, a browser would get bogged down interpreting the many thousands of tags it includes. The SVG polyline is the perfect fix for this problem: using one dramatically shrinks the size of the file and, even more, the time required to process it, since instead of a complete line tag, with its various attributes, all that is needed for each point is an <x, y> pair. Example 11-16 shows some of the code that implements this approach. Figure 11-15, which follows the pages of code, shows the result.



It might help you to understand the code in the two SVG examples if you pay more attention to its output and the way it looks in your browser than to the Python code. The Python code isn't all that intriguing: it just does what it needs to do. What's interesting is how a structured graphical image can be constructed from a structured text format.

Example 11-16. Drawing a sequence trace using SVG polylines

```
colors = ('black', 'green', 'red', 'blue')
                                                        # or whatever
axiscolor = '#333'
axisthickness = 2
tickcolor = '#666'
tickspacing = 100
                                                          # spacing of x-axis labels
tickheight = 18
tickthickness = 2
maxval = 1600
leftmargin = 10
topmargin = 20
yzero = 100
hscale = 5/2
                                                          # multiply x coordinates by this
vscale = 14
                                                          # divide y values by this
span = 1
                                                          # number of points to average together
def read data(infilname):
    with open(infilname) as infil:
        return [eval(line) for line in infil.readlines()]
def write_text(fil, x, y, txt, cls):
    print(" <text class='{}' x='{}' y='{}'>{}</text>".
          format(cls, x+leftmargin, y+topmargin, txt),
          file=fil)
def write_line(fil, frompos, fromval, topos, toval, color, thickness):
    print(" <line x1='{}' y1='{}' x2='{}' y2='{}'".</pre>
               format(leftmargin + int(frompos * hscale),
                      int(((maxval - fromval) / vscale)) + topmargin,
                      leftmargin + int(topos * hscale),
                      int(((maxval - toval) / vscale)) + topmargin),
               end=' '
               file=fil),
        print("style='stroke: {}; stroke-width: {};'/>".
               format(color, thickness),
               file=fil)
def write axes(fil, count):
    print(file=fil)
    write line(fil, 0, 0, 0, maxval, axiscolor, axisthickness)
    write line(fil, 0, 0, count, 0, axiscolor, axisthickness)
    print(file=fil)
    tickbottom = vscale * (-axisthickness - tickheight)
    for x in range(int(tickspacing / hscale),
                    count,
                    int(tickspacing // hscale)):
        write line(fil, x, tickbottom + vscale * (1 + tickheight), x, tickbottom,
                    tickcolor, tickthickness)
        write_text(fil,
                    int(x * hscale),
                    int((maxval / vscale)) + tickheight +1 5,
```

```
int(x * hscale),
                    'tick')
def write bases(outfil, start, bases, positions):
    for base, pos in zip(bases, positions):
        write text(outfil,
                    int(pos * hscale - 3),
                    int((maxval / vscale)) + 15,
                    base,
                                  # text
                    base)
                                  # class
    # - to "center" the base at its position; just a guess
    print(file=outfil)
def print_point(outfil, x, y):
    print("{},{}".format(leftmargin + int(x * hscale),
                             int((maxval - y) / vscale) + topmargin),
          file=outfil,
          end=' '
def write curve(outfil, vals, color):
    # at a point in the curve instead of using every data point
    # hscale is how far apart the x points are
    print("""
  <polyline style='fill: none; stroke: {}; stroke-width: 1;'</pre>
            points=""".
           format(color),
          file=outfil,
          end=''),
    # span is a file-level parameter that specifies the number of points to average to compute
    # a value rather than having a value for every point on the x axis; blur does the averaging
    for pos in range(0, len(vals)-1, span):
        if not pos % 6:
                                                                      # 5-digit x's at the end
            print(20*' ', file=outfil, end='')
        print point(outfil, pos/span, blur(vals[pos:pos+span]))
    print(" ' />", file=outfil)
                                                                      # close polyline
def write curves(outfil, data,):
    for d, clr in zip(data, colors):
        write curve(outfil, d, clr)
def blur(seq):
    """return the average of the values in seq"""
    return 0 if not seq else sum(seq)/len(seq)
def write heading(outfil, width, height):
    print(
"""<?xml version="1.0" standalone="no"?>
<!DOCTYPE svg PUBLIC "-//W3C//DTD SVG 1.1//EN"
"http://www.w3.org/Graphics/SVG/1.1/DTD/svg11.dtd">
```

```
<svg xmlns='http://www.w3.org/2000/svg' version='1.1'</pre>
    width='{0}' height='{1}'
>""".format(width+(2*leftmargin), height+yzero+20),
  <defs>
    <style type='text/css'><![CDATA[
        text {
              font-family: Futura, 'Andale Mono', Verdana, sans-serif;
              fill:black;
              font-weight: normal;
              font-size: 8pt;
              font-style: normal;
              text-rendering:optimizeLegibility;
            }
        text.tick { }
        text.T { fill: red; }
        text.C { fill: blue; }
        text.A { fill: green; }
        text.G { fill: black; }
    </style>
  </defs>
""", file = outfil, end='\n')
def write closing(outfil):
    print('\n</svg>', file = outfil)
def write svg file(infilname, outfilname):
    data = read data(infilname)
    start = data[0]
    bases = data[1]
    positions = data[2]
    values = data[3:]
    with open(outfilname, 'w') as outfil:
        write heading(outfil, len(values[0]) // span, maxval // vscale)
        write axes(outfil, len(values[0]) // span)
        write bases(outfil, start, bases, positions)
        write curves(outfil, values)
        write closing(outfil)
write_svg_file('data/ABI/seqdata', 'data/ABI/abi-trace.xml')
```

Figure 11-15. A portion of the generated SVG trace display

G C A A G С

A G G C A C

A G

Tips, Traps, and Tracebacks

Tips

- The first time you use a graphics library such as tkinter or a language such as SVG, the first program you write should be something very minimal. Before starting any real work, make sure that you know how to create a diagram, add a shape, display the diagram, and close it.
- Before doing any programming, sketch a high-level layout on paper. Draw vertical and horizontal lines at each significant location in your sketch. Then, invent labels for the vertical and horizontal distances in each segment of the resulting grid. Use these labels in the code you write. Figure 11-4 shows an example of what a sketch like this layout might look like, but do it by hand.
- In any kind of programming it is best to avoid using any unnamed significant numbers, but in graphics programming it is essential. Graphics code is full of numbers and computations for positions, dimensions, scaling factors, font sizes, border widths, and so on. This is the "domain" of graphics programs. You cannot avoid the numbers, but you can avoid a great deal of effort, frustration, and errors by naming every value that goes into the computation of positions and dimensions, if for no other reason than that your names will all be self-documenting.
 - If you decide to change the left margin from 12 to 18 but leave the right margin at 12, how will you know which 12s were for the right margin and which the left? Don't write code with a comment near the 12 that says "left margin." That's what names are for! In object-oriented programming class fields are the way to give these values names, as demonstrated in the programs in the first half of this chapter. For example, the histogram examples in this chapter even had a name for the number of pixels to use—3—for each 1/10 of a percent in the data. It's all right to add or subtract 1 or divide by 2 or 3 without giving the results names, but make sure the significance of those small values is trivial, not something that is controlling the nature of the graphics being constructed.
- As you develop the parts of the drawing, give each one a colored outline or background (perhaps even a different color for each one). Do this also for aspects of a region's border—that is, use different colors for padding, the border line itself, and its outside margin, depending on which of these are supported by the graphics software you are using. Coloring like this will help you detect positions and dimensions that are slightly "off" and line up pieces of the drawing.
- You don't have to develop the real parts of the drawing right away. Put in lines, rectangles, or circles that represent the positions and dimensions of the pieces you intend to add eventually. These are the graphics equivalent of pass statements.
- Concentrate on getting one part working at a time. Most programs are complex because of the interdependence of their components. Graphics programming adds

the further complication that the components of its *output* are similarly interdependent. In this chapter's examples we've seen titles, keys, axes, tics, tic labels, margins, and other elements that surround the content part of the graphics. The positions and dimensions of each of these may be affected by the positions and dimensions of several others. By programming just one component at a time you will not only avoid getting overwhelmed by the complexity of your program, but you will progressively develop a better understanding of the interrelationships among the graphical components.

- Programs that draw structured graphics can almost always be organized in several levels, as shown in this chapter's examples. High-level code simply organizes and sequences the construction of the diagram. Lower-level code typically just draws a single line or piece of text, etc. The real work is done in the middle-layer code that is called from the high-level code. The purpose of the lower-level code is to extract the repetitive patterns of method calls and parameters in the middle-level code. This can simplify the middle-level code considerably. In addition, as we saw with the tkinter-based classes in this chapter's examples, organizing your code like this will reveal similarities among the various graphics programs you write.
- You will often be able to derive a new program from an existing one with relatively few changes. Once you start doing that, you can capture the generalities in an abstract superclass and make subclasses that define only the differences from that class. The first few subclasses you develop will lead to many changes in all the classes, but after a few the superclass will generally stabilize, and you'll be able to use it easily.
- It is more important in graphics programming than in almost any other kind of software development to keep multiple versions of your files. You don't need anything fancy, just a way to save a file with a numerical suffix in a different directory. Every time you make a significant change, you should save the file both to its normal location and with an incremented backup suffix. For example, if you have five backup versions of diagram.py and have just reworked the code to take into account the border widths of some components, save it both to diagram.py and to save/diagram06.py. (Use the zero in version numbers less than 10 so that file listings are sorted appropriately.)

Traps

• "Off by one" errors and issues are pervasive in graphics programming. Avoiding them is partly a matter of learning how the tools you are using behave. For instance, a horizontal line of width 3 from (0, 10) to (50, 10) may cover the y coordinates 7-9, 8-10, 9-11, 10-12, or 11-13, depending on the software used and perhaps on parameters specified when the line is drawn. Much of the problem, though, is caused by the interdependence of component positions and dimensions. Graphics code contains a lot of computations of positions and dimensions where each

- value is a combination of several others, and often you will need to add 1 to or subtract 1 from some of these values to get things lined up properly.
- The same concerns apply to the borders of rectangles, along with some additional complications. Do their width and height values include the padding? The border? Half of the border? You'll need to become familiar with the peculiarities of your software.
- Intricacy is unavoidable in graphics code. It is rarely possible to achieve the kind of elegant terseness one should strive for in most other kinds of programming tasks. You will see just by looking at graphics code that it has an entirely different "shape" or "feel" than other code you have written or read.

SVG traps

- As in HTML, a sequence of one or more whitespace characters is treated as a single space.
- The value of each attribute must be quoted.
- Attributes are separated by spaces, not commas, semicolons, or any other punctuation.
- Style properties are *name:value* pairs separated by semicolons; whitespace may appear after the colon.
- All but text tags must be closed with a slash (i.e., />); text tags are closed with >, then followed by text and a closing </text>.
- Font properties, stroke, stroke width, fill, etc., are style properties, not tag attributes; they must be specified as the value of a tag's style attribute or in the stylesheet for the document.
- SVG's rotation operation is confusing: be sure to specify the second and third arguments for rotate transformations, or the results will almost definitely not be what you expect. In fact, rotating without an origin will usually make the element disappear from the diagram. In general, you'll want to rotate elements around either their starting points or their centers.
- CSS has a concept of "class," which is used to specify style details for all tags with a particular value for the class attribute. However, remember that class is a keyword in Python, used to define classes. Don't try to use it for a variable in code that deals with CSS classes; call it cls or cl or something similar instead. It would be nice if syntax errors caused by the use of a keyword were flagged as such in the Python error message, rather than the message simply pointing to the keyword, but that's not the case.

Tracebacks

Here's an error message you might encounter programming with tkinter:

tkinter.TclError: unknown color name "grue" A typical, and fairly helpful, tkinter error message.



One surprising thing about this error message is the underscore that precedes tkinter in its name. You will sometimes see single underscores at the beginning of names of modules that are interfaces to external libraries, such as Tk and SQLite3, or Python libraries that are implemented in C for one reason or another. They occasionally appear at the beginning of class names too. Don't worry about them—they are just ordinary names. (You can't ignore the underscores if you are trying to catch an exception whose class or module name begins with one, though: you have to give an except clause the exact name of the error class.)



Python Language Summary

This appendix summarizes the Python language as used in this book. Some of its more obscure features have been omitted.

Language Components

Special Syntactic Elements

```
# Begins a comment; the rest of the line is ignored

At end of line, indicates that line break should be ignored

"or ""

Enclose a one-line string

"" or """ """

Enclose a multiline string

()

Enclose an expression or call a function
```

Keywords

Table A-1 shows the complete list of Python's keywords. Note that a few of them play multiple roles:

- not is a logical operator and part of the not in and is not operators.
- in is an operator as well as part of a for statement.
- as names things in both with statements and except clauses.

Table A-1. Keywords

Kind	Keywords	Note
Value names	None True False	
Operators	not or and in	It is the \textit{words} that are keywords: not is both a Boolean operator
	is	and part of the not in and is not operators
	del	
General	pass assert	
	return yield	
Definitions	def class as	
	import from	
	global nonlocal	These two keywords affect the scope of assigned names; they
	lambda	are not used in any of the book's code examples.
Conditionals	if else elif	
Loops and iterations	while for break continue	$Also \verb"in", included with the operators, as part of a for statement$
Exception handling	try except finally	with is included here because it creates an implicit try/
	with raise	finally exception handler

Special Names

In addition to keywords, the names described in Table A-2 have special significance. An asterisk stands for any number of characters that can appear in a name.

Table A-2. Names with special significance

Names	Significance
_	In the interpreter only, refers to the result of the last evaluation
_*	Used for names not imported by from module import *
*	Used for names private to their classes
*_	Reserved for system-defined method names used for implementation of fundamental operations

Operators

Boolean operators are listed in Table A-3.

Table A-3. Boolean operators

Operator	Meaning
not	Unary: True if operand is False; False if operand is True
or	True if either operand is True; otherwise, False
and	True if both operands are True; otherwise, False

Table A-4 lists Python's arithmetic operators. Two of them can be used with sequences:

- + produces a new sequence with all the elements of each of its sequence operands.
- * takes a sequence and an integer and produces a new sequence with that number of repetitions of the original.

Otherwise, all operands must be numbers. If any of the operands is a float, the result will be a float.

Table A-4. Arithmetic operators

Operator	Meaning
+	Unary positive, binary addition, sequence concatenation
-	Unary negative, binary subtraction
*	Number multiplication, sequence repetition
**	Exponentiation
/	Division; result is always a float
//	Floor division; result <i>equals</i> an integer, but its <i>type</i> is an integer only if both arguments are integers
%	Remainder (modulo) of dividing the first operand by the second

Floor division and modulo are related values that are returned as tuples by the built-in function divmod:

```
divmod(n, m) == (n//m, n\%m)
n == ((n // m) * m) + (n % m)
```

Python's comparison operators are shown in Table A-5. The operands of equality and inequality may have different types, but two values of different types are always unequal. In general, it is an error to try to compare values of different types using one of the order comparisons.

Table A-5. Comparison operators

Operator	Meaning
<	op1 is less than op2
>	op1 is greater than op2
<=	op1 is less than or equal to op2
>=	op1 is greater than or equal to op2
==	op1 is equal to op2
!=	op1 is not equal to op2
is	op1 is identical to op2 (i.e., they are the same object)
is not	op1 is <i>not</i> identical to op2
in	op1 is in the collection op2
not in	op1 is not in the collection op2

Table A-6 lists the operators that perform bitwise operations. Their operands must be integers.

Table A-6. Bitwise operators

Operator	Meaning
<<	Shift op1 left by op2 bits
>>	Shift op1 right by op2 bits
&	Each bit is the and of the corresponding bit of the two operands
	Each bit is the <i>inclusive or</i> of the corresponding bit of the two operands
۸	Each bit is the <i>exclusive or</i> of the corresponding bit of the two operands
~	A unary operator: each bit is the <i>complement</i> of the corresponding bit in the operand

Anonymous Functions

A lambda expression creates an anonymous function:

lambda x, y: expression using x and y

A lambda takes one or more arguments. Only one expression—not statement—is allowed.

Types and Expressions

Table A-7 lists Python's primitive types. Collection types are summarized in Appendix B.



The str type may be considered a primitive for some purposes, and a collection for others.

Table A-7. Primitive types

Туре	Description
None	Special "no-value" value
bool	True or False
int	Integers (any magnitude)
float	"Floating-point" numbers
str	Unicode strings

Table A-8 shows the syntax for constructing expressions. (Sequences, of which strings are an example, are fully documented in Appendix B.)

Table A-8. Expressions

Kind of operator	Syntax	Notes
Numeric	+ - * / // % **	/ is float division; // is int division; $\%$ is modulo; and $\ast\ast$ is power
Logical	not and or	
Comparison	== != < <= >= >	
Sequence	+, in, not in	
Sequence subscription	[]	
Sequence slicing	[m:n], [m:n:k]	Omitting $\mbox{\ensuremath{m}}$ is equivalent to 0, omitting $\mbox{\ensuremath{n}}$ is equivalent to the end of the sequence
Function call	fn(), fn(arg1,)	
Method call	arg1.fn(arg2,)	

Statements

Table A-9 describes all the simple statements in Python. (The del statement is included here for completeness; it applies only to collection types, which are described in Appendix B.)

Table A-9. Simple statements

Statement	Description
name = expression	Binds name to the value of expression
name1 = name2 = = expression	Binds the names to the value of expression
pass	Do-nothing placeholder used where statements are required, e.g., in a function definition $ \label{eq:condition} $
return	Returns None from a function
return expression	Returns value of expression from a function
yield	Makes the function containing it a generator, with every execution of yield producing one value
assert expression	Raises an error if the expression is false
assert expression, value	$Raises an error if the expression is false, including {\it value} in the error message {\it val$
<pre>import module</pre>	Adds the contents of $module$ to Python and adds its name to the names pace in which the statement is executed
from module import name	Adds the contents of $module$ to Python and adds $name$ to the namespace in which the statement is executed
<pre>from module import name as alias</pre>	Adds the contents of <i>module</i> to Python and adds <i>alias</i> to the namespace in which the statement is executed, with <i>alias</i> used instead of <i>name</i>
del <i>coll</i> []	Removes the element(s) from coll (a list or dictionary) indicated by an index or subscription

Statement	Description
continue	Interrupts the current loop or iteration
break	Exits the current loop or iteration
<pre>raise ExceptionClass(arg)</pre>	Creates an instance of ExceptionClass and raises an exception
global name1, name2,	$\label{thm:must} Must be first statement of a function; indicates that assignments of the listed names will be in the global scope$
nonlocal name1, name2,	Must be first statement of a function; indicates that assignments of the listed names will be in the immediately enclosing scope; used when one function is defined inside of another

Table A-10 lists Python's augmented assignment statements. Both name and expression must have numeric values. (The += operator also works when the value of name is a list, in which case the value of expression may have any type; see Appendix B.) The statements on the lefthand side of the table rebind *name* to the new value, with the same effects as the statement on the righthand side of the same row of the table.

Table A-10. Augmented numeric assignment statements

Statement	Equivalent
name += expression	name = name + expression
name -= expression	name = name - expression
name *= expression	name = name * expression
name /= expression	name = name / expression
name//=expression	name = name // expression
name %= expression	name = name % expression
name **= expression	name = name * expression

There are a very small number of compound statements in Python that define new names. They are shown in Table A-11.

Table A-11. Compound statements that define names

Statement	Description
<pre>def name(parameter-list): statements</pre>	Binds name to a new function whose definition is specified by <code>parameter-list</code> and <code>body</code> , which consists of one or more statements indented relative to the word def.
class Name: statements	Defines a class, with statements one or more statements indented relative to the word class; the statements are usually defs, occasionally assignments, and very rarely anything else.
<pre>class Name(SuperName): statements</pre>	Defines a class with SuperName as its superclass.
with expression as name: statements	 Evaluates the expression. Binds <i>name</i> to the result.

Statement	Description
	3. Executes the <i>statements</i> .
	 Whether an error occurs or not, performs special actions to undo the effect of evaluating the expression.
	Used most frequently with expression, a call to open.

Table A-12 describes Python's conditional statements.

Table A-12. Conditional statements

Statement	Description
<pre>if expression: statements</pre>	Evaluates <i>expression</i> and, if true, executes the statements
<pre>if expression: statements1 else: statements2</pre>	Evaluates <i>expression</i> and, if true, executes <i>statements1</i> ; otherwise, execute <i>statements2</i>
if expression1: statements1 elif expression2 statements2	Evaluates each expression in sequence until one is true, then executes the corresponding statements; if none is true, executes <code>default-statements</code>
else: default-statements	

Table A-13 describes Python's loop and iteration statements. See Appendix B for summaries of common uses with collections.

Table A-13. Loop and iteration statements

Statement	Description
while expression: statements	Repeatedly evaluates the <i>expression</i> and executes the <i>statements</i> until the <i>expression</i> 's value is false
while expression: statements1 else: final-statements	Repeatedly evaluates the <i>expression</i> and executes the <i>statements</i> until the <i>expression</i> 's value is false; then evaluates <i>final-statements</i>
<pre>for item in collection: statements</pre>	For each item in collection, executes the statements

An exception statement begins with the keyword try and must have at least one of the other kinds of clauses. There can be any number of except clauses that list one or more exception classes. There may not be more than one except clause that does not name an exception class, nor more than one finally clause.

When an exception is raised during the execution of the try block (that is not handled by a "deeper" try statement), that block is immediately exited. Next, the type of the exception raised is compared to the type(s) of each except clause until a match is found, at which point the statements of that clause are executed and execution continues after the last except clause. If an except clause without any classes is reached, its statements are executed.

Note that it is possible for all the except clauses to list one or more class names, none of which match the type of the exception raised. In that case, the try statement does not catch the exception, and Python continues looking up the stack for a statement that does catch the exception.

Regardless of whether the try statements completed without an exception being raised or were interrupted by an exception, if there is a finally clause, its statements will be executed.

The components of an exception handling statement are shown in Table A-14. Note that what appears in the lefthand side of the statement are *clauses*, not *statements*. A try statement must try and have one or more except clauses. It may have one finally clause.

Table A-14. Exception handling statements

Statement	Description
try: try-statements	Executes the try-statements
<pre>except ExceptionClass[,]: except-statements</pre>	If an exception that is an instance of one of the listed types is raised, executes the <code>except-statements</code> ; then continues after the last <code>except</code> clause of the <code>try</code> statement
<pre>except ExceptionClass[,] as name: except-statements</pre>	As in the form described in the previous row, but also names the exception object so it can be referenced in the <code>except-statements</code>
finally: finally-statements	Executes the <i>finally-statements</i> regardless of whether an error occurred or not; must be the last in the series of clauses that follow the <i>try-statements</i>

Notes

Two features of Python syntax were omitted from this book because they disturb the regularity of code appearance. Since you may see these in other code you read, though, you should know about them:

1. Strings can be concatenated during compilation, as opposed to execution, simply by putting them next to each other with no separating +. They can even appear on separate lines, as long as all that is between them is whitespace. For example:

```
print('This string might be too long to fit on one line where'
       it appears in your program, so just write two or more'
      ' strings in succession')
```

There's no particular advantage to doing this rather than using a +, except in the rare case where a statement containing a concatenation of string literals (not names whose values are strings) would otherwise be executed a very large number of

- times. Concatenation at compile time creates a single string. Concatenation at runtime creates a new string every time a string concatenation expression (*str1* + *str2*) is evaluated.
- 2. When the body of a one-clause compound statement (or the body of a clause of a multiclause compound statement) consists of a single statement, it can appear on the same line as the first line of the clause, after the colon. For example:

```
if value: print(value)
else: print('None found')
```



Collection Type Summary

This appendix provides a compact summary of (most of) Python's collection types. It describes the characteristics and methods of each collection and the operators and functions used with collection values. At the end of the appendix, you will find summaries of the collection iteration templates shown in Chapter 4.

Types and General Operations

Table B-1 summarizes the characteristics of Python's most commonly used collection types. The column labeled "Syntax" shows examples of two-element collections, with e1 and e2 representing elements of any type and e3 representing instances of an immutable type.

Table B-1. Collection types

Element access	Name	Туре	Element type	Mutable?	Syntax
Unordered unique	Set	set	Any immutable	Yes	$\{k1, k2\}$
elements	Frozenset	frozenset	Any immutable	No	None
Ordered (indexed)	String	str	One-character	No	'ac' "ac"
			strings ^a		'''ac''' """ac"""
	Bytes	bytes	8-bit bytes ^b	No	b'ac' b"ac"
					b'''ac''' b"""ac"""
	Bytearray	bytearray	8-bit bytes	Yes	None
	Range	range	Integers	No	None
	Tuple	tuple	Any	No	(e1, e2)
	List	list	Any	Yes	[e1, e2]
Associative	Dictionary	dict	Keys immutable, values any	Yes	{k1:e1,

Element access	Name	Туре	Element type	Mutable?	Syntax
					k2:e2}
Stream (next)	File object	Depends ^c	Characters, bytes, lines	Depends on use	None

^a Strings don't really "contain" characters or one-character strings, but for many purposes they behave as if they do.

Common Operations

For all sequence types except range, calling the type with no arguments produces an empty sequence, and calling the type with a sequence argument produces a new object of the type containing the elements of the argument.

The two byte types—bytes and bytearray—can be called with an integer to create a sequence of that many zero bytes.

When dict is called with a sequence argument the elements of the sequence must all be sequences of length 2 containing a key and a value, in that order.

Except for range, all sequence types implement the methods count and index. Table B-2 lists the operations and functions that apply to all collection types.

Table B-2. Operations and functions common to all collection types

Function/operation	Returns
x in coll	True if coll contains x.
x not in <i>coll</i>	True if <i>coll</i> does not contain <i>x</i> .
any(coll)	True if any item in <i>coll</i> is true; otherwise, False.
all(coll)	True if every item in <i>coll</i> is true; otherwise, False.
len(coll)	The number of items in <i>coll</i> .
max(coll)	The maximum item in coll, which may not be empty.
min(coll)	The minimum item in coll, which may not be empty.
sorted(coll[, keyfn] [, reverseflag])	A list containing the elements of <i>coll</i> , sorted by comparing elements or, if <i>keyfn</i> is included in the call, comparing the results of calling <i>keyfn</i> for each element; if <i>reverseflag</i> is true, the ordering is reversed. <i>keyfn</i> and <i>reverseflg</i> must be specified as keyword arguments, not positionally.

Creating Collections

As with all types, collection types can be called to create new instances. Table B-3 shows the details of the parameters of each type's call.

b Bytes are equivalent to the integers from 0 through 255.

The type of the object returned by open is an instance of the class from the IO library; it isn't one of the built-in collection types.



In the Python documentation, you will find these functions in the section "Built-in Functions," under "Built-in Types."

Table B-3. Collection creation calls

Method	Description	
<pre>set([iterable])</pre>		
<pre>frozenset([iterable])</pre>		
<pre>str([source[, encoding]])</pre>	If no <i>source</i> , returns an empty string; if no <i>encoding</i> , returns the result of <i>source</i> 'sstr method if it has one, orrepr if not. If the type of <i>source</i> is bytes or bytearray, decodes it using <i>encoding</i> .	
<pre>bytes([source[,</pre>	If <i>source</i> is an integer, returns a bytes instance of that many null bytes; an iterable of integers 0<=x<=255 is used to initialize the new bytearray. If <i>source</i> is a string, <i>encoding</i> must be specified (e.g., 'utf8') and will be used to encode <i>source</i> .	
<pre>bytearray([source[, encoding]])</pre>	Same as with bytes.	
<pre>range([start], stop[, step])</pre>	Arguments must be integers; returns a range object from $start$ (default 0) to $stop$ in increments of $step$ (default 1).	
<pre>tuple([iterable])</pre>	Returns a tuple with the items of iterable in the order in which they appear there.	
<pre>list([iterable])</pre>	Returns a list with the items of $iterable$ in the order in which they appear there.	
<pre>dict([arg])</pre>	With no axg , returns an empty dictionary; with axg a dictionary, returns a copy. Otherwise, axg must be an iterable of key/value pairs.	
open(filename, mode)	Creates a file object.	

Another way to create collections is with comprehensions:

```
List comprehension:
[expression for item in collection]
Set comprehension:
{expression for item in collection}
Dictionary comprehension:
{key-expression: value-expression
for key, value in collection}
```

The collection in a dictionary comprehension must consist of two-element tuples or lists.

Conditions can be added to comprehensions. There can be any number of tests, and all have to be true for the element to be included in the result. Here are some examples for lists:

```
[expression for item in collection if test]
[expression for item in collection if test1 if test2]
```

Comprehensions can have more than one for. For example:

```
[c1 + c2 + c3 for c1 in 'TCAG' for c2 in 'TCAG' for c3 in 'TCAG']
```

Specific Collection Types

This section summarizes the nature of the collection types used in this book and the functions, methods, and expressions that operate on them.

Sets

A set is an unordered collection of items that contains no duplicates.

Table B-4 lists comparison operations that can be performed on sets. Some are expressed using an operator whose operands are both sets, some with a method call whose argument is any kind of collection, and some with either.

Table B-4. Set comparison operations

Operator	Method	Result
	<pre>set1.isdisjoint(coll)</pre>	True if the set and the argument have no elements in common
set1 <= set2	<pre>set1.issubset(coll)</pre>	True if every element of set1 is also in set2 (col1 for the method)
set1 < set2		True if every element of set1 is also in set2 (col1 for the method) and set2 is larger than set1 (i.e., set1 is a proper subset of set2)
set1>= set2	<pre>set1.issuperset(coll)</pre>	True if every element of set 2 (col1 for the method) is also in set 1
set1>set2		True if every element of set2 (coll for the method) is also in set1 and set1 is larger than set2 (i.e., set2 is a proper subset of set1)

Table B-5 lists operations that produce new sets from the elements of other sets or collections. Each can be expressed using an operator whose operands are both sets. Three of them have an equivalent method whose arguments are any number of any kind of collection. One has an equivalent method that takes just one collection of any type.

Table B-5. Algebraic set operations

Operator	Method	Result
set1 set2	set1.union(coll,)	A new set with the elements of both the set and the arguments
set1&set2	<pre>set1.intersection(coll,)</pre>	A new set with the elements that are common to the set and the arguments

Operator	Method	Result
set1 - set2	<pre>set1.difference(coll,)</pre>	A new set with the elements that are in the set but not in the arguments
set1 ^ set2	<pre>set1.symmetric_difference(coll)</pre>	A new set with the elements that are in either the set or the arguments but not both

Table B-6 lists the operations and methods that actually modify the first set rather than returning a new set.

Table B-6. Set modification operations

Operator	Method	Result
set1 = set2	<pre>set1.update(coll)</pre>	Updates <i>set1</i> by adding the elements in <i>set2</i> (<i>col1</i> for the method)
set1&= set2	<pre>set1.intersection_update(coll)</pre>	Updates set1 to keep only the elements that are in both set1 and set2 (col1 for the method)
set1 - set2	<pre>set1.difference_update(coll)</pre>	Updates $set1$ to keep only the elements that are in $set1$ but not in $set2$ ($col1$ for the method)
set1^set2	<pre>set1.symmetric_differ ence_update(coll)</pre>	Updates set1 to keep only the elements that are in either set1 or set2 (col1 for the method)
	<pre>set1.add(item)</pre>	Adds item to set1
	<pre>set1.remove(item)</pre>	Removes <i>item</i> from <i>set1</i> ; it is an error if <i>item</i> is not in the set
	<pre>set1.discard(item)</pre>	Removes item from set1 if it is present; no error if it is not present

Sequences

Sequences are ordered collections that may contain duplicate elements.

Table B-7 summarizes sequence indexing and slicing operations.

Table B-7. Sequence indexing and slicing

Type of seq	Indexing	Slicing	Slice assignment
	seq[n]	seq[i:j]	seq[i:j] =obj
str	Yes	Yes	No
bytes	Yes	Yes	bytes no
bytearray			bytearray yes
range	Yes	No	No
tuple	Yes	Yes	No
list	Yes	Yes	Yes

Table B-8 summarizes other sequencing operations.

Table B-8. Sequence operators

Type of seq	Membership	Combination	Repetition	Comparison*
	obj in seq	seq1 + seq2	seq*n	$seg\cdot other$
	obj not in seq	(seq1 and seq2 have the same type)	(nanint)	with · one of ==, ! =, >, <, >=, or <=
str	<i>obj</i> is a str	str	Yes	other is a str
bytes	<i>obj</i> is 0255, a	Yes	Yes	other is a bytes or
bytearray	bytes,ora bytearray			bytearray
range	<i>obj</i> is an int ^a	No	No	No
tuple	obj has any type	Yes	Yes	other is a tuple
list	obj has any type	Yes	Yes	other is a list

^a Actually, the expressions obj in xng and obj not in xng are always valid for any type of obj and any range xng, but always False if

Table B-9 documents the way various forms of slice expressions work.

Table B-9. Slice expressions

Operation	Returns
seq[i:j]	Elements of seq from $m{i}$ up to, but not including, $m{j}$
seq[i:]	Elements of seq from i through the end of the sequence
seq[:j]	Elements of seq from first up to, but not including, j
seq[:-1]	Elements of seq from first up to, but not including, the last
seq[:]	All the elements of seq—i.e., a copy of seq
seq[i:j:k]	Every $k^{ ext{th}}$ element of s eq , from $m{i}$ up to , but not including, $m{j}$

Strings

Strings are sequences of Unicode characters (though there is no "character" type). The bytes and bytearray types are sequences of single bytes. They have essentially the same operations and methods as str. One important difference is that with a string, regardless of whether you index an element or specify a slice, you always get back a string. With bytes and bytearrays, although slices return the same type, indexing an element returns an integer from 0 through 255.

Since strings and bytes are immutable, no str or byte method modifies the string through which it is called. However, bytearray is a mutable type; elements of its instances may be replaced by using assignment, including assignment of slices.

Table B-10 lists string predicate methods. All predicates return False for an empty string. For the "case" tests, the string must contain at least one character of the appropriate type. Note that not all characters are "cased": digits and punctuation, for example, are not.

Table B-10. String predicates for kinds of characters

Method	Result
isalpha	True if all characters are alphabetic
isalnum	True if all characters are alphanumeric
isdigit	True if all characters are digits
isnumeric	True if all characters are numeric, including Unicode number characters
isdecimal	True if all characters are valid for use in decimal numbers
islower	True if all cased characters are lowercase
isupper	True if all cased characters are uppercase
istitle	${\bf True}\ if all\ upper case\ characters\ follow\ uncased\ characters\ and\ all\ lower case\ characters\ follow\ cased\ characters$

Table B-11 describes the string methods that search for one string inside another. In all the methods documented, startpos defaults to 0 and endpos defaults to the end of the string (inclusive)—i.e., str2 is compared to str1, str1[startpos:], str1[:endpos], or str1[startpos:endpos], according to which of the optional arguments are specified in a call.

Table B-11. String predicate and search methods

Mashad	Decula
Method	Result
str1.startswith(str2[, startpos[, endpos]])	True if $str1$ starts with $str2$, taking into account $startpos$ and $endpos$
str1.endswith(str2[, startpos[, endpos]])	True if $str1$ ends with $str2$, taking into account $startpos$ and $endpos$
str1.find(str2[, startpos[, endpos]])	Lowest index of $str2$ in $str1$, taking into account $startpos$ and $endpos; -1$ if not found
str1.rfind(str2[, startpos[, endpos]])	Highest index of $str2$ in $str1$, taking into account $startpos$ and $endpos; -1$ if not found
str1.index(str2[, startpos[, endpos]])	Lowest index of $str2$ in $str1$, taking into account $startpos$ and $endpos$; error if not found
str1.rindex(str2[, startpos[, endpos]])	Highest index of $str2$ in $str1$, taking into account $startpos$ and $endpos$; error if not found
str1.count(str2[, startpos[, endpos]])	The number of occurrences of $str2$ in $str1$, taking into account $startpos$ and $endpos$

Table B-12 lists methods that return a new string with characters adjusted as necessary to the corresponding case.

Table B-12. String methods for changing case

Method	Result
str1.lower()	A copy of $str1$ with all of its characters converted to lowercase
str1.upper()	A copy of $str1$ with all of its characters converted to uppercase
<pre>str1.capitalize()</pre>	A copy of $str1$ with only its first character capitalized; has no effect if the first character is not a letter (e.g., if it is a space)
str1.title()	A copy of $str1$ with each word beginning with an uppercase character and the rest lowercase
str1.swapcase()	A copy of $str1$ with lowercase characters made uppercase and vice versa

Methods that remove spaces from or add them to strings are described in Table B-13.

Table B-13. Methods for reformatting strings

Method	Result
<pre>str1.lstrip([chars])</pre>	A copy of str1 with all characters until the first character not contained in chars (default is whitespace) removed
<pre>str1.rstrip([chars])</pre>	Like lstrip, but removes characters from the end
<pre>str1.strip([chars])</pre>	Like lstrip, but removes characters from both the start and the end
str1.ljust(width[, fillchar])	A string that is at least $width$ long, padded with $fill char$ on the left (default is whitespace)
str1.rjust(width[, fillchar])	Like ljust, but the padding is on the right
<pre>str1.center(width[,</pre>	Like ljust, but the padding is divided between the left and right
<pre>str1.center([tabsize])</pre>	A string with each tab of $str1$ replaced with enough spaces to reach the next multiple of $tabsize$ (default is 8)

Methods for joining and splitting strings are documented in Table B-14.

Table B-14. Methods for joining and splitting strings

Method	Result
sepr.join(seq)	A string formed by concatenating the strings in the sequence seq , separated by $sepx$, which can be any string (including the empty string)
<pre>str1.splitlines([keepflg])</pre>	A list of the lines in $str1$; if $keepf1g$ is True, the string includes end-of-line characters
<pre>str1.split([sepr [,</pre>	A list of the "words" in str1, using sepr as a word delimiter (default is whitespace); if count is included as an argument, the result is limited to count words

Method	Result
<pre>str1.rsplit([sepr [,</pre>	Like split, but starts from the end
<pre>str1.partition(sepr)</pre>	A tuple of three elements: the portion of $str1$ up to the first occurrence of $sepr$, $sepr$, and the portion of $str1$ after the first occurrence of $sepr$; if $sepr$ is not found, the result is $(s,'','')$
str1.rpartition(sepr)	Like partition, but starts from the end

Table B-15 lists some general-purpose string replacement methods.

Table B-15. General-purpose string replacement methods

Method	Result
<pre>str1.replace(oldstr,</pre>	Returns a copy of str1 with all occurrences of the substring oldstr replaced by the string newstr; if count is specified, only the first count occurrences are replaced.
<pre>str1.translate(dictionary)</pre>	With dictionary having integers as keys, returns a copy of str1 with any character char for which ord(char) is a key in dictionary replaced by the corresponding value; exactly what the replacement does depends on the type of the value in the dictionary, as follows:
	 None—character is removed from str1
	 Integer n—character is replaced by chr(n)
	 String str2—character is replaced by str2, which may be of any length
<pre>str.maketrans(x[, y[, z]])</pre>	(Called directly through the str type, not an individual string.) Produces a translation table for use with translate more conveniently than manually constructing the table; arguments are interpreted differently depending on how many there are:
	 x—character is removed from str1
	 x, y—x is a dictionary like that expected by translate, except that its keys may be either integers or one-character strings
	 x, y, z—x and y are strings of equal length; the table will translate each character of x to the character in the corresponding position of y

Lists

Lists can be modified through assignment statements, as shown in Table B-16.

Table B-16. List-modifying assignments

Statement	Result
lst[n] = x	Replaces the n^{th} element of 1st with x
lst[i:j] = coll	Replaces the \emph{i}^{th} through \emph{j}^{th} elements of \emph{lst} with the elements of $\emph{col1}$

Statement	Result
<pre>lst[i:j] = empty_coll</pre>	Deletes the i^{th} through j^{th} elements of lst (an important special case of $lst[i:j] = seq$)
<pre>lst[i:j:k] = coll</pre>	Replaces the elements of lst designated by the slice with the elements of $coll$, whose length must equal the number of elements designated by the slice
<pre>lst[n:n] = coll</pre>	Inserts the elements of $coll$ before the n^{th} element of lst
lst[len(lst):len(lst)] = [x]	Adds x to the end of $1st$
lst[len(lst):len(lst)] = coll	Adds the elements of $coll$ at the end of lst
lst+=coll	
lst[:] = coll	Replaces the entire contents of $1st$ with the elements of $col1$

The methods that modify lists are shown in Table B-17.

Table B-17. List-modifying methods

Method	Result
<pre>lst.append(x)</pre>	Adds x to the end of $1st$
<pre>1st.extend(x)</pre>	Adds the elements of x at the end of $1st$
<pre>lst.insert(i, x)</pre>	Inserts x before the i^{th} element of $1st$
<pre>lst.remove(x)</pre>	Removes the first occurrence of x from $1st$; raises an error if x is not in $1st$
<pre>1st.pop([i])</pre>	Removes the i^{th} element from $1st$ and returns it; if i is not specified, removes the last element
<pre>lst.reverse()</pre>	Reverses the list
<pre>lst.sort([reverseflag] [, keyfn])</pre>	Sorts the list by comparing elements or, if <i>keyfn</i> is included in the call, comparing the results of calling <i>keyfn</i> for each element; if <i>reverse flag</i> is true, the ordering is reversed; <i>keyfn</i> and <i>reverseflg</i> must be specified as keyword arguments, not positionally

Mappings

A mapping is a mutable unordered collection of key/value pairs. The only mapping type used in this book is dict. Expressions and statements for dictionaries are documented in Table B-18.

Table B-18. Dictionary expressions and statements

Operation	Result
d[key]	Returns the value associated with key ; raises an error if d does not contain key
d[key] = value	Associates <i>value</i> with <i>key</i> , either adding a new key/value pair or, if <i>key</i> was already in the dictionary, replacing its value

Operation	Result
d[key] ·= value	Augmented assignment, with \cdot any of $+$, $-$, $*$, $/$, $/$, $\%$, or $**$; raises an error if d does not have a value for key or if the value is not numeric
del <i>d</i> [<i>key</i>]	Deletes <i>key</i> from the dictionary; raises an error if <i>d</i> does not contain <i>key</i>

Table B-19 lists methods that are unique to dict objects.

Table B-19. Methods unique to dictionaries

Method	Result
<pre>d.get(key[, default])</pre>	Like $d[key]$, but does not cause an error if d does not contain key ; instead it returns $default$, which, if not provided, is None
d.setdefault(key[, default])	Like d[key] if key is in d; otherwise, adds key with a value of default to the dictionary and returns default (if not specified, default is None)
<pre>d.pop(key[,default])</pre>	Like $del\ d[key]$, but does not cause an error if d does not contain key ; instead it returns $default$, which, if not provided, is None
<pre>d1.update(d2)</pre>	For each key in $d2$, sets $d1[key]$ to $d2[key]$, replacing the existing value if there was one
d.keys()	Returns a special sequence-like object containing the dictionary's keys
<pre>d.values()</pre>	Returns a special sequence-like object containing the dictionary's values
<pre>d.items()</pre>	Returns a special sequence-like object containing $(key, value)$ tuples for the dictionary's keys

Streams

A stream is a temporally ordered sequence of indefinite length, usually limited to one type of element. Each stream has two ends: a source that provides the elements and a *sink* that absorbs the elements.

File objects

File objects are created by calling open(filename, mode). Tables B-20 and B-21 show the characters that can be included in the *mode* string.

Table B-20. Mode values for opening a file—text or binary

	Mode	Interpretation
t	Text (default)	Characters or strings depending on method
b	Binary	Bytes

Table B-21. Mode values for opening a file—use

	Initial file position	Read?	Write?
r	Beginning (default)	Yes	No
W	Beginning	No	Yes
a	End	No	Yes
r+	Beginning	Yes	No
W+	Beginning	Yes	Yes
a+	End	Yes	Yes

Table B-22 describes the methods supported by file objects. Note that the read methods in this table return empty strings when called at the end of the file, and that they may be called repeatedly at the end of the file with no error occurring. Also, although writelines is named analogously to readlines, it does not add end-of-file characters to the lines it writes; if you want them, you need to include them explicitly in the lines in the sequence. Another way to write to a file is to specify it as the value of the optional file argument to the function print.

Table B-22. File object methods

Method	Result
<pre>file.read([count])</pre>	Reads and returns $count$ bytes or to the end of $file$, whichever comes first; if $count$ is omitted, reads to end of file
<pre>file.readline([count])</pre>	Reads and returns $count$ lines or to the end of $file$, whichever comes first; if $count$ is omitted, reads to end of file
<pre>file.readlines()</pre>	Reads and returns lines until the end of $file$ is encountered
<pre>file.write(str)</pre>	Writes str to file
<pre>file.writelines(seq)</pre>	Writes the strings contained in the sequence seq to file

Generators

A generator is a stream that returns values from a series it computes. The only thing you can do with a generator is call next on it to get its next value:

next(generator[, default])

Gets the next value from the *generator* object. If *generator* has no more values to produce, returns default if specified in the call; otherwise, raises the StopIteration exception.

Generators may be created in one of two ways:

• A function definition that uses yield in place of return returns a generator object when it is called.

• The value of an expression with syntax like a comprehension, except with parentheses instead of brackets or braces, is a generator.

Iteration Templates

Most iterations conform to a limited set of patterns. The outlines in this section's tables use the following abbreviations:

```
-item-
    Any use of item
?-item-?
    Boolean expression using item
-item1, item2-
    Any use of two items together
    Any operator (or function call)
fn
    Any function name
```

Basic Iteration Templates

The iteration templates are outlined in Table B-23.

Table B-23. Basic iteration templates

Section	Action	Code outline
"Do" on page 114	Do	<pre>for item in collection: -item-</pre>
"Collect" on page 115	Collect (comprehension usually better)	<pre>result = [] for item in collection: result.append(-item-) return result</pre>
"Combine" on page 117	Combine	<pre>result = initial_value for item in collection: result = result · -item- return result</pre>
"Combine" on page 117	Count	<pre>count = 0 for item in collection: count += 1 return count</pre>
"Combine" on page 117	Collection Combine	<pre>result = [] for item in collection: result += fn(item) return result</pre>

Filtering Templates

Templates in which operations are applied only to elements meeting criteria expressed in some Boolean test are summarized in Table B-24.

Table B-24. Filtering templates

Section	Action	Code outline
"Search" on page 120	Search	<pre>for item in collection: if ?-item-?: return item</pre>
"Filter" on page 122	Filtered Do	<pre>for item in collection: if ?-item-?: -item- return result</pre>
"Filter" on page 122	Filtered Collect	<pre>result = [] for item in collection:</pre>
	(conditional comprehension usually better)	<pre>if ?-item-?: result.append(-item-) return result</pre>
"Filter" on page 122	Filtered Combine	<pre>result = initial_value for item in collection: if ?-item-?: result = result · -item- return result</pre>
"Filter" on page 122	Filtered Count	<pre>count = 0 for item in collection: if ?-item-?: count += 1</pre>
		return count

Other Kinds of Templates

A few other templates are described in Table B-25.

Table B-25. Other templates

Section	Action	Code outline
"Nested iterations" on page 126	Nested Do (with similar variations for other kinds of templates)	<pre>for item1 in collection1: for item2 in collection2: -item1,item2-</pre>
"Recursive itera- tions" on page 128	Recursive Iteration	A function definition that implements an iteration during which the function calls itself (or calls another function that calls it, etc.); needed for recursive structures

Index

Symbols != (comparison operator), 8 # symbol, 2, 28 % (operator), 5 * (binary operator), 9 ** (power operator), 5 + (binary operator), 9 / (operator), 5	dot plots example, 416 examples, 4, 16 FASTA files, 77–78 generating translations of codons, 86 amino acid translation table, 68–71 and (operator) Boolean logic, 7 seach loops, 109
// (operator), 5 // (operator), 5 < (comparison operator), 8 <= (comparison operator), 8 == (comparison operator), 8 > (comparison operator), 8 >= (comparison operator), 8 @ notation, 182 character, 265	anonymous functions, 92–94, 452 any (function) collection arguments, 48, 111 dictionary support, 71 lambda expressions and, 93 arc (as graphic component), 403 arguments defined, 12 evaluating in functions, 13 functions as, 91, 92
A a (HTML tag), 291–294 abs (function) defined, 14 parameter considerations, 89	lambda expressions, 92 in method calls, 15 methods as, 91 parameters and, 27 sequences as, 61
passing values, 91 absolute URLs, 332 access methods, 168–170 action attribute (HTML), 350	specifying for function calls, 13 arithmetic operators augmented assignment statements and, 23 overview, 5–7
action methods, 177 all (function) collection arguments, 48, 111 dictionary support, 71 lambda expressions and, 93 list comprehensions, 80	assertion statements defined, 31 overview, 30–32 representative error messages, 46 with two-expressions, 31 AssertionError (error), 46, 137
ALTER TABLE (SQL statement), 371, 377 amino acid sequences	assignment statement bindings and, 27

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bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 Canvas (class) create_arc (method), 408 create_oval (method), 408 create_polygon (method), 408 create_rectangle (method), 408	class attributes and, 179	assigning default values, 32
comprehension results and, 111 defined, 23 associative arrays (see mappings) AttributeError (error) classes, 208 control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 creation calls, 54 defined, 73 bytes, defined, 73 bytes, defined, 73 bytes, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 poperators supporting, 7–9 predicates and, 48 supported values, 2 bound methods, 208, 316 boundaries, 208, 316 boundaries, regular expressions, 261 brace notation, 263 break statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytea, defined, 21 creation calls, 54 defined, 21 creation ca	class method and, 183	comparison operators, 8
defined, 23 associative arrays (see mappings) AttributeError (error) classes, 208 control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating random, 81 base, numeric defined, 27 translating random, 81 base, numeric defined, 27 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 23 defined, 21 exception statements, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 predicates and, 48 supported values, 2 bound methods, 208, 316 boundaries, regular expressions, 261 brace notation, 26 brace notation, 26 brace notation, 263 break statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes(type) built-in for sequences, 52 creation calls, 54 defined, 73 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 21 exception statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements of the formation, 48 lists and, 63 bytes (type) built-in for sequences of the formation, 48 lists and, 63 bytes (type) built-in for sequences of the formation, 48 lists and, 63 bytes (type) built-in for sequences of the format	classes as objects, 184	operator precedence order and, 17
defined, 23 associative arrays (see mappings) AttributeError (error) classes, 208 control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating random, 81 base, numeric defined, 23 in exidence, 53 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 predicates and, 48 supported values, 2 bound methods, 208, 316 boundaries, regular expressions, 261 brace notation, 26 brace notation, 26 brace notation, 26 brace notation, 263 break statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bottom controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences of the defined, 21 creation calls, 54 defined, 73 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequence of the defined, 21 creation	comprehension results and, 111	operators supporting, 7–9
associative arrays (see mappings) AttributeError (error) classes, 208 control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 139 function parameters, 27 with statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 supported values, 2 bound methods, 208, 316 boundaries, regular expressions, 261 brace notation, 263 boundaries, regular expressions, 261 brace notation, 26	defined, 23	
AttributeError (error) classes, 208 control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bitts, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 bound methods, 208, 316 boundaries, regular expressions, 261 brace notation, 26 brace notation, 26 brace notation, 26 break statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes, defined, 73 bytes attament iteration support, 112 loop interruption and, 104, 105 buffers example, 278–282 flushing, 278 button controls, 351 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes arangle attaments, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes arangle, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes and, 63 bytes (type) built-i		
classes, 208 control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bitms, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 boundaries, regular expressions, 261 brace notation, 26 brace notation, 26 brace notation, 26 brace statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 32, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (cass) create_line (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408		
control statements, 163 example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating random, 81 base, numeric defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 break statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearrayy (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences of sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54	classes, 208	
example, 137 attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 break statement iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 crea		
attributes class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 iteration support, 112 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 21 call sacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method		
class, 179–185 defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 loop interruption and, 104, 105 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 23 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54		
defined, 167 structured text and, 287 augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 buffers defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 33 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 63 bytes (type) built-in for sequences, 52 creation calls, 64 defined,		
structured text and, 287 augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 defined, 73, 278 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 25 function parameters, 27 with statements, 139 function parameters, 27 function parameters, 27 with statements, 139 function parameters, 27 functio		
augmented assignment statements, 23, 62 B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 2 function calls to, 14 lambda expressions and, 93 example, 278–282 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 40 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 40 defined, 53 bytes (type) built-in for sequences portanting of the fined particular for the fined particu		
B backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 2 function calls to, 14 lambda expressions and, 93 flushing, 278 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytearrays in assignment statements, 63 lists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes, defined, 73 call stacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_oval (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408		
backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 2 function calls to, 14 lambda expressions and, 93 button controls, 351 bytearray (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 2 feration calls, 54 defined, 3 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 2 feration calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 2 feration calls, 54 defined, 2 feration calls of, 4 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 3 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 3 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 3 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 3 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 3 bytes (type) built-in for sequences, 52 creation calls, 64 defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas	augmented assignment statements, 25, 62	
backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 2 function calls to, 14 lambda expressions and, 93 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (typ	_	
backslashes, 258, 260 base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 2 freation calls, 54 defined, 2 freation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls (s) bytes (type) call season, 63 bytes (type) creation calls, 54 defined, 53 bytes (type) call season, 63 bytes (type) call season, 63 bytes (type) creation calls (s) defined, 2 functio	В	
base classes, defined, 194 base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 bitmaps, 399 create_inc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408	backslashes, 258, 260	
base sequences reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 cetattont calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) creation calls, 54 defined, 53 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes (type) calls facks, 310–311 callbacks, 311–314 calls calls defined, 53 bytes (type) calls facks (also, 10–31 calls facks, 310–311 callbacks, 311–314 calls calls defined, 53 bytes (type) calls f		
reading from FASTA files, 146–147 translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 call stacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408		
translating example, 86, 151–154 translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes, defined, 73 bytes, 310–311 callbacks, 310–311 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408		
translating random, 81 base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 iists and, 63 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 call stacks, 310–311 callbacks, 311–314 calls oelined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408		
base, numeric defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 bytes (type) built-in for sequences, 52 creation calls, 54 defined, 53 bytes, defined, 73 bytes, defined, 73 call stacks, 310–311 callbacks, 311–314 calls create, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408		
defined, xxii hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bytes, defined, 73 Call stacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408 create_rectangle (method), 408 create_rectangle (method), 408	_	*
hexidecimal notation, 2 binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 butter not sequences, 32 creation calls, 54 defined, 53 bytes, defined, 73 call stacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
binary operators defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 call stacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
defined, 5 operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bool (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 bytes, defined, 73 bytes, defined, 73 call stacks, 310–311 callbacks, 311–314 calls defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_oval (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
operator precedence order and, 17 string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bool (HTML tag), 288, 296 bool (type) defined, 2 function and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		defined, 53
string operations, 9–12 binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bool (HTML tag), 288, 296 bool (type) defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		bytes, defined, 73
binding assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 booly (HTML tag), 288, 296 bool (type) defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
assignment statements, 23 defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bool (HTML tag), 288, 296 bool (type) defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		(
defined, 21 exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 bool (HTML tag), 288, 296 bool (type) defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		-
exception statements, 139 function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
function parameters, 27 with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function (alls to, 14 lambda expressions and, 93 defined, 12 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
with statements, 76 bioinformatics additional information, xxi main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function, 12–15 method, 15–16 operator precedence order and, 17 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_polygon (method), 408 create_rectangle (method), 408 create_rectangle (method), 408		
bioinformatics method, 15–16 additional information, xxi operator precedence order and, 17 main areas, xii canvas software development, xii overview, 408 bitmaps, 399 writing contents to files, 410 bits, defined, 73 canvas (class) body (HTML tag), 288, 296 create_arc (method), 408 bool (type) create_line (method), 408 defined, 2 create_oval (method), 408 function calls to, 14 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408		
additional information, xxi operator precedence order and, 17 main areas, xii Canvas overview, 408 writing contents to files, 410 bits, defined, 73 Canvas (class) body (HTML tag), 288, 296 create_arc (method), 408 bool (type) create_line (method), 408 create_oval (method), 408 function calls to, 14 create_polygon (method), 408 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408		
main areas, xii software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 Canvas overview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_rectangle (method), 408		
software development, xii bitmaps, 399 bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 cverview, 408 writing contents to files, 410 Canvas (class) create_arc (method), 408 create_line (method), 408 create_oval (method), 408 create_polygon (method), 408 create_rectangle (method), 408		operator precedence order and, 17
bitmaps, 399 writing contents to files, 410 bits, defined, 73 Canvas (class) body (HTML tag), 288, 296 create_arc (method), 408 bool (type) create_line (method), 408 defined, 2 create_oval (method), 408 function calls to, 14 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408		Canvas
bits, defined, 73 body (HTML tag), 288, 296 bool (type) defined, 2 function calls to, 14 lambda expressions and, 93 Canvas (class) create_arc (method), 408 create_oval (method), 408 create_polygon (method), 408 create_rectangle (method), 408	software development, xii	overview, 408
body (HTML tag), 288, 296 create_arc (method), 408 bool (type) create_line (method), 408 defined, 2 create_oval (method), 408 function calls to, 14 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408	bitmaps, 399	writing contents to files, 410
bool (type) create_line (method), 408 defined, 2 create_oval (method), 408 function calls to, 14 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408	bits, defined, 73	Canvas (class)
defined, 2 create_oval (method), 408 function calls to, 14 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408	body (HTML tag), 288, 296	create_arc (method), 408
function calls to, 14 create_polygon (method), 408 lambda expressions and, 93 create_rectangle (method), 408	bool (type)	create_line (method), 408
lambda expressions and, 93 create_rectangle (method), 408	defined, 2	create_oval (method), 408
lambda expressions and, 93 create_rectangle (method), 408		
Boolean logic, 2 create_text (method), 408	Boolean logic, 2	create_text (method), 408
(see also bool (type)) Cascading Style Sheets (CSS), 393		Cascading Style Sheets (CSS), 393

CGI (Common Gateway Interface)	overview, 166–168
defined, 343	relationships with methods, 186-205
serving requests, 343	representative error messages, 208
setting up, 344	storing values, 168
cgi (module), 343	tips using, 205
CGI scripts	traps using, 207
arguments and responses, 345–347	types and, 1
debugging, 346	clauses
HTML forms and, 350–354	defined, 100
PyChart library support, 411	headers in, 100
simple applications, 348–350	suites as synonym, 100
cgi.FieldStorage (class), 346	coding considerations
cgi.test (function), 345	event-based processing, 314–317
CGIHTTPRequestHandler (class), 343, 350	infinite loops, 107
character classes (in regular expressions), 261	tips developing and testing, 95
character sets (in regular expressions), 260	tips executing, 41
circle (as graphic component), 403	collections
class attributes	built-in type categories, 47
accessing, 179	comprehensions, 79–89
class fields, 179–182	comprehensions and, 117
class methods, 182-184	creating, 47, 460
classes as objects, 184-185	functional parameters, 89-94
overview, 179	items or elements, 47
typical uses, 179	mappings, 66–72
class decomposition, 186–189	objects and, 47, 111
class definitions	purposes of, 167
decomposing, 186	representative error messages, 97
defined, 166	sequences, 51–66
statement template, 167	sets, 48–51
class fields	streams, 72–79
common uses, 180	tips using, 94
overview, 179–182	traps using, 96
tracking instance counts, 183	types and general operations, 459–462
class methods	colon
assignment statements and, 183	in class definition statement, 167
defined, 182	in compound statements, 24, 99
distinguishing from instance methods, 182	dictionaries and, 67
overview, 182–184	format specifier, 59
Tracking Instance Count template, 183	in headers, 100
classes	command-line interpreter
basic template, 178	editing operations supported, 19
class attributes, 179–185	quitting, xx
defined, xxiii	running Python interactively, 18
determining which to define, 167	streams into, 73
inheritance, 194–205	command-line utilities
instance attributes, 168–178	fileinput (module), 217
namespaces and, 179	optparse (module), 218–220
as objects, 184–185	subprocess (module), 221–223
objects as instances, 1	comments

# symbol in, 2, 28	containers (see collections)
defined, 2	continue statement
docstrings versus, 29	iteration support, 112
overview, 28–30	loop interruption and, 104, 105
COMMIT (SQL statement), 373	control statements
Common Gateway Interface (see CGI)	conditionals, 101-103
communications	defined, 100
logging (module), 225	exception handlers, 134-142
smtplib (module), 224	extended examples, 143–160
comparison operators	iterations, 111–132
defined, 8	loops, 104–110
operator precedence order and, 17	representative error messages, 163
for sets, 49	tips using, 160–162
composite (as graphic component), 403	traps using, 162
compound expressions, 16–17	coordinate system, structured graphics, 400-
compound statements	402
defined, 24, 99	CREATE TABLE (SQL statement), 371, 375
headers in, 100	CSS (Cascading Style Sheets), 393
kinds listed, 101	csv (module), 241
compound types (see collections)	csv.reader (function), 241
comprehensions	csv.writer (class)
collections and, 117	writerow (method), 242
conditional, 87–88	writerows (method), 242
conditional iterations, 111	csv.writer (function), 242
defined, 79	cubic spline (as graphic component), 403
dictionary, 85	curly braces
generator expressions and, 85	creating sets, 48
list, 80–84	dictionaries and, 67
nested, 88, 111	empty, 48
set, 85	format specifiers, 58
concatenating strings, 9	
conditional comprehensions, 87–88	D
conditional expressions	-
defined, 8	database servers, 359
loops with guard conditions, 109	databases (see relational databases)
conditional statements	datetime (module)
defined, 101	classes supported, 209
infinite loops, 107	instance creation, 210
multi-test, 102	methods supported, 212
one-alternative, 102	operations supported, 211
simple, 101	overview, 209
connection objects	datetime.date (class), 210 isocalendar (method), 212
abort (method), 373	isoweekday (method), 212
close (method), 373	replace (method), 211
commit (method), 373	today (class method), 210
creating, 372–374	datetime.datetime (class), 210
execute (method), 373	combine (class method), 210
executemany (method), 373	date (method), 211
executescript (method), 373	isocalendar (method), 212
	isocarciluai (ilictilou), 212

isoformat (method), 212	setdefault (method), 72
isoweekday (method), 212	update (method), 72
now (class method), 210	values (method), 72, 98, 112, 163
replace (method), 211	dict type, 67
time (method), 211	dictionaries
datetime.time (class), 210	empty, 48
isoformat (method), 212	indexing and removal, 71
replace (method), 211	iteration support, 112
datetime.timedelta (class), 210	methods supported, 72
datetime.tzinfo (class), 210	operations supported, 71, 113
dbm (module), 243–246	overview, 67–68
dbm.dumb (module), 245	RNA codon translation table, 68–71
dbm.gnu (module), 245	sets comparison, 67
firstkey (method), 245	dictionary comprehensions
nextkey (method), 245	conditional, 88
reorganize (function), 245	defined, 85
dbm.ndbm (module), 244	difflib (module), 237
dbm.open (function), 244	difflib.Differ (class), 237
debugging	compare (method), 238
CGI scripts, 346	difflib.HtmlDiff (class), 238
IDLE debugger, 252	make_file (method), 238
overview, 249	make_table (method), 238
pdb (module), 250–252	difflib.SequenceMatcher (class), 237
regular expressions, 283	find_longest_match (method), 237
decomposition	get_matching_blocks (method), 237
class, 186–189	set_seq1 (method), 237
defined, 186	set_seq2 (method), 237
method, 189–193	set_seqs (method), 237
singleton classes, 193	dir (function), 63, 87
def statement	directories
defined, 24, 91	comparing with files, 235–238
defining methods, 167	listing contents, 271
del statement	managing, 227
defined, 64	distributed computing, 248
dictionary support, 72	division operators, 5–7
delegation, defined, 189	divmod (function), 61
DELETE (SQL statement), 371	do-nothing statement (pass), 26
deleting elements, 64	docstrings
derived classes, defined, 194	comments versus, 29
dict	defined, 29
defined, 72	Document Type Definition language, 300
function call errors, 98	documenting functions, 28–30
iterations and, 112	dot notation, 35, 167
representative error messages, 163	dot plots
dict (type), 67	additional information, 416
get (method), 72	defined, 399, 416
items (method), 72, 98, 112, 163	graphics example, 416–423
keys (method), 72, 98, 113, 163	DROP TABLE (SQL statement), 371, 376
pop (method), 72	, <u> </u>

E	statements supporting, 138–141
	tracebacks, 136
Element (class), 304	exception raising
find (method), 305	ending loops, 142
findall (method), 305	extracting HTML file information, 143
findtext (method), 306	146
get (method), 305	raise statement, 141
getchildren (method), 306	expat (module) (see xml.parsers.expat
getiterator (method), 306	(module))
items (method), 305	expressions
keys (method), 305	compound, 16–17
set (method), 305	comprehension results in, 111
values (method), 305	defined, xxii, 5
elements, collection	function calls, 12–15, 91
defined, 47	generator, 85–87
deleting, 64	lambda, 92–94, 316, 452
elements, structured text, 287	logical operators, 7–9
ElementTree	method calls, 15–16
navigation, 305–309	numeric operators, 5–7
overview, 303–305	Python language summary, 452
ElementTree (module)	statements and, 23
find (method), 305	string operations, 9–12
findall (method), 305	tips using, 18
findtext (method), 306	
getiterator (method), 306	two-expression assertions, 31
getroot (method), 306	Extensible Markup Language (see XML) extension, defined, 204
parse (function), 304	extension, defined, 204
elif clause, 102	<u>_</u>
else clause	F
in conditionals (if statements), 102	factoring out common code, 198-199
in loops (while statements), 104	FASTA files
else keyword (conditional expression), 8	Collect iteration example, 116
email messages, sending, 224	collecting information example, 134
end tags (HTML), 288	Combine iteration example, 118
Entrez Gene site, 143	extracting entries, 274–282
enumerate (function), 113	Filtered Do iteration example, 123
environment variables, 234	loops with guard conditions, 110
EOFError (error), 137, 163	reading, 77–78, 82–84, 87
event loops, 406	reading sequences from, 146–147
event-based processing	Search iteration example, 120–122
callbacks, 311–314	fields (see class fields)
function calls, 310–311	FieldStorage (class)
programming considerations, 314–317	getfirst (method), 346
except clause (try statement), 138, 139	getlist (method), 346
Exception (class), 142	_
	file keyword, 77
exception handlers	file objects
event-based processing and, 310–311	creating, 74
generators and, 141	iteration support, 112
overview, 134	methods supported, 76
runtime errors, 136–138	read (method), 76

readline (method), 76, 78, 86, 110	finally clause (try statement), 139
readlines (method), 76, 86	fixed-length matching
summarized, 469	boundaries, 261
write (method), 76	character classes, 261
writelines (method), 76	character sets, 260
filecmp (module), 235–237	literal matches, 259
cmp (function), 235	flag parameters, 32
cmpfiles (function), 235	flat files, defined, 287
filecmp.dircmp (class), 236	float (type)
report (method), 236	defined, 3
report_full_closure (method), 236	function calls to, 14
report_partial_closure (method), 236	floating point numbers
fileinput (module), 217	decimal points and, 3
filelineno (function), 217	integer division and, 5
filename (function), 217	scientific notation and, 3
isfirstline (function), 217	floor division, 5
lineno (function), 217	flushing buffers, 278
nextfile (function), 218	fnmatch (module), 233
files	filter (function), 233
comparing with directories, 235–238	fnmatch (function), 233
defined, 73	for statement
flat, 287	basic form, 111
functionality in streams, 73–74	Collect template, 115–116
importing, 38	Collection Combine template, 119
as interfaces, 73	Combine template, 117–119
listing contents of directories, 271	Count template, 118
looping over lines in, 110	dictionary iteration, 112
managing, 227	Do template, 114
manipulating contents, 77–78	file iteration, 112
parsing, 147	Filtered Collect template, 123–125
showing differences between, 237	Filtered Combine template, 125–126
as stream sinks, 73	Filtered Count template, 126
as stream sources, 73	Filtered Do template, 122
temporary, 229	nested comprehensions, 88, 111
tips managing, 44	Nested Iteration template, 126–128
filesystem utilities	numbering iterations, 113
difflib (module), 237	Recursive Tree Iteration template, 128–132
filecmp (module), 235–237	Repeat template, 115
fnmatch (module), 233	Search template, 120–122
glob (module), 233	form (HTML tag), 350
os (module), 226–228	format (function), 58–60
os.path (module), 229–232	format specifiers
shutil (module), 234	defined, 58
tempfile (module), 229	general fields, 59
filesystems, defined, 128	numeric fields, 59
filtering	numeric type fields, 60
conditional comprehensions and, 87	forward slash, 332
defined, 87	frames (HTML)
iteration support, 122–126	defined, 310, 405
neration support, 122–120	defilled, 510, 705

event-based processing, 310	class decomposition example, 187–189
querying databases, 392–395	file parsing example, 148–150
frameset (HTML tag), 393	method decomposition example, 189-193
frozenset (type)	singleton classes example, 193
defined, 48	generalization of subclasses, 200-203
mathematical operations, 49	generator expressions
function calls	conditional, 88
callbacks and, 311–314	defined, 85
comprehension results in, 111	overview, 85–87
event-based processing and, 310–311	generators
as expressions, 91	defined, 73, 78
flow of control and, 100	exception handling and, 141
operator precedence order and, 17	loop considerations, 108
overview, 12–15	as stream sources, 73
specifying arguments, 13	summarized, 470
to types, 14	yielding values from, 79
function definitions	geometry managers, 405
defined, 24	glob (module), 233
overview, 24–34	glob (function), 234, 338, 340
tips using, 40	iglob (function), 234
function names, 12	Grand Unified Bioinformatics File Parser, 147
function parameters	graphics programming
arguments and, 27	building full-featured class, 411–416
default values, 32–34	concepts, 400–403
key parameter, 89–91	coordinate system, 400–402
overview, 27–28	GUI toolkits, 404–406
function return (see generators; return	image components, 402
statement)	overview, 399
functions	property considerations, 403
	greedy matching, 263
anonymous, 92–94, 452 as arguments, 91, 92	GTK+ toolkit, 405
	guard conditions
assertion statements, 30–32	
built-in, 13–15	defined, 109
classes as objects, 184	loops with, 109–110
collection arguments, 48	GUI toolkits
comments in, 28–30	overview, 404
components, 12	for Python, 404
decomposing, 186	usage considerations, 405
defined, 12	
documentation in, 28–30	Н
generators as, 73	hash tables (see mappings)
initializing loop values, 106	head (HTML tag), 296
method decomposition, 189	headers, defined, 100
passing values to, 13, 24, 91	heading (HTML tag), 295–296
recursive, 129	help (function)
representative error messages, 46	defined, 14
	looking at docstrings, 29
G	hexadecimal notation, 2
GenBank	histograms
	0

www.it-ebooks.info

defined, 399	dictionary support, 71, 113
graphics example, 424–430, 436	sequence types, 53
href (HTML attribute), 291–294, 325	string operations, 9
HTML	IndentationError (error), 163
CGI scripts and, 350-354	indenting code
constructing/viewing pages, 330	blank lines and, 28
elaborate pattern matching, 295–296	tab characters and, 24
extracted links and, 333	IndexError (error), 20, 98, 137
extracting information example, 143-146	indexes
nested tags, 298	accessing sequences, 47
problems with pattern matching, 297	defined, 9
querying databases, 392–395	dictionary support, 71
searching text, 290–294	negative values, 10
structured processing, 297–299	slicing strings, 10–12
structured text and, 287–289	infinite loops
turning into plain text, 296	coding, 107
html.parser (module), 298–299	defined, 105
handle_comment (method), 299	information hiding principle, 166
handle_data (method), 299	inheritance
handle_endtag (method), 298, 314	defined, 194
handle_startendtag (method), 299	defining subclasses, 195–197
handle_starttag (method), 298	factoring out common code, 198–199
HTMLParser (class), 298, 314	generalization of subclasses, 200–203
http.server (module), 341–342	multiple, 195
•	subclass methods, 203–205
1	init (method)
<u> </u>	class fields example, 180
IDLE	class methods example, 183
debugging support, 252	defining subclasses and, 195
tips using, 41–43	method decomposition and, 193
traps using, 207	overview, 172
if keyword (conditional expression), 8	as superclass extension, 204
if statement	validating arguments, 198
conditional iteration, 111	initialization methods, 172
multi-test conditional form, 102	input (function), 13
one-alternative conditional form, 102	input controls, 351
simple condition form, 101	INSERT INTO (SQL statement), 371, 378
img (HTML tag), 291	instance dictionary, setting up, 180
immutable types, 47	instance fields
import statement	defined, 167
overview, 34–38	determining which to access, 169
Python files and, 38	initialization, 172–173
selective, 35	modification methods for, 175
ImportError (error), 46, 254	storing values, 168
importing	instance methods
files, 38	access methods, 168-170
modules, 34–38	action methods, 177
representative error messages, 46	distinguishing from class methods, 182
in (operator)	initialization methods, 172
collection arguments, 48, 111	

modification methods, 175–177	K
predicate methods, 171–172	
string methods, 173–175	KeyboardInterrupt (error), 137, 163
support methods, 177	KeyError (error), 137, 163
instances	keys
defined, 1	accessing mappings, 47
testing value types, 15	defined, 360
tracking, 180	keywords
int (type)	defined, 27
defined, 2	in headers, 100
function calls to, 14	parameter properties, 33
integers	Python language summary, 449
dividing, 5	
hexadecimal notation, 2	L
repeating strings, 9	-
string operations, 9	lambda expressions, 92–94, 316, 452
interfaces, files as, 73	LANG environment variable, 295
	layout managers, 405
interpreter (see command-line interpreter)	leaves, defined, 129
io (module), 242	len (function)
io.ByteIO (class), 242	collection arguments, 48, 111
io.StringIO (class), 242	defined, 13
IOError (error), 137, 163	Filtered Count iteration and, 126
isinstance (function), 15, 125	key parameter, 94
items (see elements)	li (HTML tag), 288
iterations	libraries
Collect template, 115–116	defined, xxiii
Collection Combine template, 119	modules and, 34
Combine template, 117–119	line (as graphic component), 402
Count template, 118	list (type), 52
defined, 111	append (method), 64
dictionaries, 112	extend (method), 64, 65
Do iteration, 114	insert (method), 64
file objects, 112	pop (method), 64
Filtered Collect template, 123–125	remove (method), 64
Filtered Combine template, 125–126	reverse (method), 64
Filtered Count template, 126, 306	sort (method)
Filtered Do template, 122	defined, 64
filtering support, 122–126	key parameter, 89
for statement support, 111–113	lambda expressions and, 94
kinds supported, 113–132	predicate method example, 171
Nested Iteration template, 126–128	list comprehensions
numbering, 113	conditional, 87
overview, 111	defined, 80–84
Recursive Tree Iteration template, 128–132,	lists
308	in assignment statements, 63
Search template, 120–122	bytearrays and, 63
template summary, 471–472	defined, 62
	modification assignment expressions, 63
	modification methods, 64
	· · · · · · · · · · · · · · · · · · ·

slicing, 63	collection arguments, 48, 111
suffix trees and, 130	defined, 14
summarized, 467	dictionary support, 71
literal matches, 259	key parameter, 89
logging (module), 225	measurement, units of, 401
logical operators	method calls
conditional expressions, 8	operator precedence order and, 17
operator precedence order and, 17	overview, 15–16
overview, 7–9	method decomposition, 189–193
lookup tables (see mappings)	methods, 182
loops	(see also class methods; instance methods)
basic form, 104	as arguments, 91
defined, xxiii, 104	association with types, 22
exception raising and, 142	defined, 15
with final clause, 104	dictionary, 72
generators and, 108	file object, 73, 76
with guard conditions, 109–110	instance attributes, 168–178
infinite, 105, 107	list, 64
initializing values, 106–107	referring to superclasses, 195
interrupting, 104	relationships with classes, 186–205
over lines of files, 110	sequence, 65
search, 109	sets, 49, 50, 51
simple examples, 105	string, 15
lt (method)	subclass, 203–205
class fields example, 180	methods, bound, 208
class methods example, 183	min (function)
overview, 171	collection arguments, 48, 111
representative error messages, 208	defined, 14
	dictionary support, 71
M	key parameter, 89
many-many relationships, 362, 389	modification methods, 175–177 modules
mappings	
accessing elements of, 47	defined, 34
as collection type, 47	filesystem, 226–238
dictionaries, 67–72	importing, 34–38
summarized, 468–469	namespaces and, 35–37
unique keys, 67	persistent storage, 243–252
match objects	Python files and, 38
defined, 258, 265	representative error messages, 46, 255
findall (method), 268	system environment, 209–226
finditer (method), 268	tips using, 253
match (method), 268	traps using, 254
re (module), 269–270	working with text, 238–243
search (method), 268	modulo, defined, 5
split (method), 268	multiple inheritance, 195
sub (method), 268	
subn (method), 268	N
matching (see pattern matching)	name attribute (HTML), 350
max (function)	named entities, 288
max (runction)	manieu chitics, 200

NameError (error)	files as interfaces, 73
classes, 208	as instances of classes, 1
collections, 97	persistent, 247–248
example, 137	values and, 1
undefined names, 46	ol (HTML tag), 288
names	one-many relationships, 362, 387
assigning to values, 23	one-one relationships, 362
conflict considerations, 36	open (function), 74
defined, 21	operands
function, 12	Boolean operators, 8
strings versus, 21	defined, 5
tips using, 40	string operations, 9
namespaces	operators
association with types, 22	compound expresssions, 16–17
classes and, 179	defined, 5
defined, 21	logical, 7–9
function parameters and, 27	numeric, 5–7
modules and, 35–37	precedence rules, 16
nested comprehensions, 88, 111	Python language summary, 450–452
nested HTML tags, 298	string operations, 9–12
nested iterations, 126–128	optparse (module), 218–220
network connections as stream sources, 73	or (operator), 7
network sources as stream sinks, 73	os (module)
next statement	chdir (function), 227
defined, 79	environ (variable), 226, 347
generator expressions, 86	environment access, 226
non-greedy matching, 263	getcwd (function), 227
None value	getenv (function), 226
defined, 1	listdir (function), 227
print (function) and, 114	managing files/directories, 227–228
search iterations and, 120	mkdir (function), 227
search loops and, 109	mkdirs (function), 227
not (operator), 7	overview, 226
not in (operator)	remove (function), 227
collection arguments, 48, 111	removedirs (function), 227
dictionary support, 71	rename (function), 227
sequence types, 53	rmdir (function), 227
string operations, 9	sep (variable), 226
numeric operators	startfile (function), 227
operator precedence order and, 17	walk (function), 227
overview, 5–7	os.path (module), 229
	abspath (function), 230
^	basename (function), 230
0	dirname (function), 230
object-oriented programming, 22	exists (function), 231
objects	expanduser (function), 231
binding, 23	getsize (function), 231
classes as, 184–185	isdir (function), 231
in collections, 47, 111	isfile (function), 231
file-supported, 73	

www.it-ebooks.info

join (function), 231	polyline (as graphic component), 402
split (function), 230	PostScript files, 410
splitdrive (function), 230	power operator
splitext (function), 230	defined, 5
OSError (error), 255	operator precedence order and, 17
oval (as graphic component), 403	pprint.pprint (function)
overriding, defined, 204	dictionary example, 70, 71
OWL (Web Ontology Language), 301	iterations and, 114
	list comprehensions, 82
P	precedence rules, operators, 16
•	predicate methods, 171–172
parameters (see function parameters)	predicates, defined, 48, 171
parentheses, 195, 264	primary keys, 360
parsing defined, 147	primitives
	expressions, 5–17
GenBank file example, 148–150	as immutable types, 47
method decomposition and, 190	overview, 1
singleton classes and, 193	Python language summary, 452
XML files, 301 pass statement, 26	representative error messages, 20
	simple values, 1–5
paths, manipulating, 229–232	tips using, 18
pattern matching examples, 270–282	traps for, 20
fixed-length, 259–262	print (function)
	defined, 13
fundamental syntax, 258–265 greedy versus non-greedy, 263	None value, 114
HTML considerations, 295–296	optional argument, 77
problems with HTML, 297	string methods and, 174
re (module), 265–270	programming (see coding considerations;
regular expressions and, 257	graphics programming; web
	programming)
representative error messages, 285 searching versus, 258	programming languages
tips using, 283	criteria for clear writing, 103
traps using, 284	overview, xv-xvii
variable-length, 262	properties, graphics, 403
pdb (module), 250–252	PyChart library, 411
percent-encoding, 325	Python
period, 261	additional information, xx
persistent storage utilities	exiting, 19
dbm (module), 243–246	installing, xviii
pdb (module), 250–252	language summary, 449–456
pickle (module), 247–248	modules concept, xi
shelve (module), 248	overview, xvii
pickle (module), 247–248	running, xix–xx
dump (function), 248	running interactively, 18
dumps (function), 247	syntax considerations, xvii
load (function), 248	terminology, xxi
loads (function), 248	
policies, layout, defined, 405	Q
polygon (as graphic component), 403	Qt toolkit, 405
porygon (as grapine component), 403	Quitolikit, 105

quadratic spline (as graphic component), 403	re.IGNORECASE flag, 296
queries	re.search method, 296
constructing and submitting, 329	re.sub (method), 296
qualified, 385	readline library, 19
relationship, 386	Rebase data, 348, 365–369, 374–380
simple, 381–385	rectangle (as graphic component), 402
SQL, 380–391	recursive functions, 129
from web pages, 392–395	recursive iterations, 128–132
question marks, 387	refactoring, defined, 204
quotation marks surrounding strings, 5	reformatting strings, 57
quotation marito ourro arraing orinigo, o	regular expressions
n	additional information, 257
R	debugging, 283
raise statement, 141	defined, 258
random (module), 37	disjunction, 264
randint (function), 37, 78	extracting sequence file descriptions, 272–
random numbers	274
generating, 37	—· ·
list comprehensions and, 81	extracting sequence file entries, 274–282
range (function), 60, 80	findingopen reading frames, 271–272
range (type)	fixed-length matching, 259–262
built-in for sequences, 52	boundaries, 261
overview, 60	character classes, 261
ranges, defined, 60	character sets, 260
rational numbers, 7	literal matches, 259
raw string syntax, 258	fundamental syntax, 258–265
raw_input (function), 13	greedy versus non-greedy matching, 263
RDF (Resource Description Framework), 301	grouping, 264
_	ignoring case, 270
flore supported 266, 267	listing files in directories, 271
flags supported, 266–267	pattern matching and, 257
functions supported, 265–266	raw string syntax, 258
match objects, 269–270	repetition characters, 262
methods supported, 268–269	testing, 258
overview, 265	traps using, 284
re (module)	variable-length matching, 262
ASCII (flag), 266	relational data models, 359
compile (function), 266, 268	relational databases
DOTALL (flag), 266, 284	avoiding duplication of values, 360-362
escape (function), 266	database definition, 364
findall (function), 265	querying from web pages, 392-395
finditer (function), 266	representative error messages, 398
IGNORECASE (flag), 266, 271	representing relationships, 362–364
LOCALE (flag), 266	restriction enzyme example, 365–369
match (function), 265	tips using, 397
MULTILINE (flag), 266, 273, 284	traps using, 398
search (function), 265	relationships, representing, 362–364
split (function), 266	relative URLs, 332
sub (function), 266	RELAX NG language, 300
subn (function), 266	repetition characters, regular expressions, 262
VERBOSE (flag), 266, 273, 284	repetition characters, regular expressions, 202

REPLACE (SQL statement), 371	as arguments, 61
replacing strings, 56	built-in types, 52
repr (method)	bytearrays, 53
class fields example, 180	bytes, 53
class methods example, 183	as collection type, 47
overview, 173–175	creating strings, 54
reserved words, 27	defined, xxii, 51
Resource Description Framework (RDF), 301	formatted text output, 58-60
restriction enzymes	generally supported operations, 52
database example, 365–369	list comprehensions and, 80
literal matches, 259	lists, 62–66
return statement	overview, 51–53
comprehension results in, 111	ranges, 60
loop interruption and, 105	reformatting strings, 57
overview, 13, 24	replacing strings, 56
search iterations and, 120	searching strings, 55
traps using, 45	slicing, 52
tuple packing and, 61	string methods, 65
reversed (function), 53	summarized, 52, 463–468
rows attribute (HTML), 393	testing strings, 55
runtime errors, 136–138	tuples, 61–62
	servers (see web servers)
S	set (type), 48, 49
	add (method), 51
Scalable Vector Graphics (see SVG)	difference (method), 50
scientific notation, 3	difference_update (method), 51
script (HTML tag), 296	discard (method), 51
searching	intersection (method), 50
HTML text, 290–294	intersection_update (method), 51
iteration support, 120–122	isdisjoint (method), 49
loops with guard conditions and, 109	issubset (method), 49
matching versus, 258	issuperset (method), 49
strings, 55	remove (method), 51
SELECT (SQL statement)	symmetric_difference (method), 50
defined, 371	symmetric_difference_update (method), 51
DISTINCT keyword, 382	union (method), 50
expressions in, 382	update (method), 51
LIMIT clause, 381	set comprehensions, 85
qualified queries, 385	sets (type)
simple queries, 381–385	accessing elements of, 47
WHERE clause, 385	algebraic operations, 49
selective import, 35	as collection type, 47
Semantic Web, 301	comparison operators, 49
sequence (see amino acid sequences; base	creating, 48
sequences)	defined, 48
sequence files	dictionary comparison, 67
extracting descriptions, 272–274	frozenset (type), 48
extracting entries, 274–282	mathematical operations, 49
sequences types	overview, 48–51

summarized, 462	(see also specific types of statements)
update operations, 50	examples of simple, 99
shape tags (SVG), 433	modifying lists, 63
shelve (module), 248	Python language summary, 453–456
open (function), 249	tips using, 18
shutil (module), 234	values and, 23
Simple Mail Transport Protocol (SMTP), 224,	StopIteration (error)
337	event-based processing, 315, 318
singleton classes, 193	generators and, 87, 141, 142
slicing	storage (see persistent storage utilities)
defined, 9, 10–12	str (type)
lists, 63	built-in for sequences, 52
operator precedence order and, 17	capitalize (method), 57
sequences, 52	center (method), 58
SMTP (Simple Mail Transport Protocol), 224,	commonly used methods, 15
337	count (method), 15, 56
smtplib (module), 224	creation calls, 54
smtplib.SMTPAuthenticationError (error),	defined, 4, 53
255	endswith (method), 55
smtplib.SMTPConnectError (error), 255	expandtabs (method), 58
socket.error message, 255, 342, 358	find (method), 15, 55, 142
socket.send (function), 340	format (method), 58
socket.sendall (function), 340	function calls to, 14
sockets	index (method), 56
defined, 337	isalnum (method), 55
web servers and, 337–342	isalpha (method), 55
sorted (function)	isdecimal (method), 55
collection arguments, 48	isdigit (method), 55
predicate methods and, 171	islower (method), 55
span (HTML tag), 295	istitle (method), 55
special methods, defined, 171	isupper (method), 55, 171
SQL (Structured Query Language)	join (method), 66
defined, 359	ljust (method), 58
major operations, 371–380	lower (method)
query support, 380–391	defined, 57
sqlite3 (module)	key parameter, 90
connecting to database, 372–374	passing values, 91
installing, 372	lstrip (method), 16, 57
overview, 371	maketrans (method), 56
relationship queries, 386	numeric (method), 55
sqlite3.Error (class), 373, 398	partition (method), 66, 82
sqlite3.OperationalError (class), 398	replace (method), 56, 296
square brackets	rfind (method), 56
enclosing optional arguments, 13	rindex (method), 56
for lists, 63	rjust (method), 58
slicing strings, 10	rpartition (method), 66
src attribute (HTML), 393	rsplit (method), 66
start tags (HTML), 288	rstrip (method), 16, 57
statements, 100	split (method), 66, 77, 83

splitlines (method), 66	testing, 55
startswith (method), 15, 55	as Unicode characters, 4
strip (method), 16, 57	structured graphics
swapcase (method), 57	defined, 399
title (method), 57	graphics programming, 399-406
translate (method), 56	representative error messages, 447
upper (method), 57	SVG support, 431–440
str (method)	tips using, 444
class fields example, 180	tkinter toolkit, 406–430
class methods example, 183	traps using, 445
overview, 173–175	Structured Query Language (see SQL)
stream sinks	structured text
common examples, 73	defined, 287
defined, 72	HTML format, 287-299
stream sources	representative error messages, 323
common examples, 73	tips using, 322
defined, 72	traps using, 323
streams	XML support, 300–321
accessing elements of, 47	subclass methods, 203-205
buffers and, 73	subclasses
defined, 47, 72	defined, 194
files and, 73–78	defining, 195–197
summarized, 469–470	generalization of, 200–203
string (module), 240–241	subprocess (module), 221–223
capwords (function), 240	subprocess.call (function), 221
string methods, 173–175	subprocess.getoutput (function), 221
strings, 4	subscription
(see also regular expressions; str (type))	defined, 9–10
accessing as collections, 47	operator precedence order and, 17
binary operators manipulating, 9–12	suffix trees, 129
changing case, 57	suites (see blocks)
compound expressions, 17	sum (function), 117
concatenating, 9	super (function), 195
creating, 54	superclasses
defined, xxii, 53	defined, 194
formatted output, 58–60	defining subclasses, 195
as immutable types, 47	factoring out common code, 198
line wrapping considerations, 5	multiple inheritance, 195
names versus, 21	subclass methods and, 203–205
quotation marks surrounding, 5	support methods, 177
reformatting, 57	SVG (Scalable Vector Graphics)
repeating, 9	additional information, 431
replacing, 56	characteristics, 431
searching, 55	defined, 431
sequence-oriented methods, 65	font properties, 435
slicing, 9, 10–12	graphics examples, 436–440
spanning multiple lines, 4	shape tags, 433
subscribing, 9–10	style attributes, 434
summarized, 464–467	style properties, 435

SVG File template, 432–435	gmtime (function), 216
SyntaxError (error), 20, 97, 398	sleep (function), 216
sys (module), 213	tkinter (module)
argv, 213	basic steps, 406
builtin_module_names, 214	Canvas drawing methods, 408-410
exit, 215, 315	documentation, 405
modules, 214	graphics examples, 411-430
path, 214	representative (error) messages, 447
stderr, 214	widget options, 407
stdin, 214	writing canvas contents to files, 410
stdout, 214, 279, 345	trace file curves
system environment utilities	defined, 399
datetime (module), 209–212	SVG example, 440
fileinput (module), 217	tracebacks, defined, 136
logging (module), 225	transactions, defined, 374
optparse (module), 218–220	translating
smtplib (module), 224	defined, xxiii
subprocess (module), 221–223	RNA sequences, 86, 151–154
sys (module), 213–215	trees, recursive iterations and, 128–132
time (module), 216	try statement
	exception handling, 138
T	exception raising, 142
	optional features, 139–141
tab characters, 24	tuple (type)
tempfile (module), 229	built-in for sequences, 52
gettempdir (function), 229	sequences as arguments, 61
makedtemp (function), 229	tuples
makestemp (function), 229	defined, 61
temporary files, 229	packing and unpacking, 61, 113
testing	syntax supported, 61
code, 95	two-expression assertion statement, 31
regular expressions, 258	type attribute (HTML), 351
simple conditional statements and, 102	TypeError (error)
strings, 55	argument errors, 46
value types, 15	classes, 208
text (as graphic component), 402	collections, 98
text files, constructing tables from, 155–160	control statements, 163
text manipulation utilities	example, 137
csv (module), 241	web programming, 358
io (module), 242	types, 1
string (module), 240–241	(see also collections; specific types)
textwrap (module), 238	calling as functions, 14
textwrap (module), 238	classes and, 1
dedent (function), 238	defined, xxii, 1
fill (function), 239	immutable, 47
wrap (function), 238	namespaces and, 22
tilde character, 234	values as instances, 1
time (module), 216	versions of methods, 22
clock (function), 216	
ctime (function), 216	

U	(see also None value)
unary operators	assigning names to, 23
defined, 5	bool (type), 2
operator precedence order and, 17	for default parameters, 32–34
underscore, 87, 171	defined, 1
Unicode characters	entering, 1
additional information, 4, 53	float (type), 3
file objects and, 74	in indexes, 10
HTML considerations, 288	initializing in loops, 106–107
	as instances of types, 1
UnicodeDecodeError (error), 323	int (type), 2
units of measurement, 401	objects and, 1
unwinding process, 311	passing to functions, 13, 24, 91
UPDATE (SQL statement), 371	statements and, 23
urllib.error.HTTPError (error), 358	storing for classes, 168
urllib.error.URLError (error), 358	str (type), 4
urllib.parse	testing types, 15
assembling URLs, 327	yielding from generators, 79
disassembling URLs, 326	variable-length matching, 262
overview, 325	vocabulary, defining, 32
urllib.parse (module)	von Willebrand disease, 143
quote (function), 327	
quote_plus (function), 327	W
unquote (function), 327	= =
unquote_plus (function), 327	web clients
urldefrag (function), 327	absolute URLs, 332
urlencode (function), 327	dowloading linked files, 334–337
urljoin (function), 327, 333	extracted links, 333
urlparse (function), 326	overview, 331
urlunparse (function), 327	Web Ontology Language (OWL), 301
urllib.request (module), 331	web programming
geturl (function), 331	manipulating URLs, 325–330
info (function), 331	representative error messages, 358
urlopen (function), 331, 333, 335	tips using, 356
urlretrieve (function), 331	traps using, 357
URLs	web clients, 331–337
absolute, 332	web servers, 337–354
assembling, 327	web servers
constructing/submitting queries, 329	CGI support, 343–354
contructing/viewing HTML pages, 330	defined, 337
disassembling, 326	http.server (module), 341–342
manipulating, 325–330	overview, 338–340
relative, 332	simple applications, 348
utility modules (see modules)	sockets and, 337–342
	webbrowser (module), 328, 331
V	open (function), 328, 329
•	open_new (function), 328
value attribute (HTML), 351	open_new_tab (function), 328
ValueError (error), 137	while statement
values, 1	basic loop form, 104

loop interruption and, 104 loop with final clause, 104 recursive structures and, 128 wildcard characters filename expansion, 232 pattern matching and, 258 with statement exception handling, 141 overview, 75 wxWidgets toolkit, 405

X

XML (Extensible Markup Language) ElementTree (module), 303–309 event-based processing, 310-317 expat (module), 317-322 genome example, 302 overview, 300-302 parsing files, 301 xml.etree.cElementTree (module), 301 xml.etree.ElementTree (module), 303-309 xml.parsers.expat (module), 301, 317-322 CharacterDataHandler (field), 318 ParseFile (method), 317 Parser (method), 317 ParserCreate (function), 317 StartElementHandler (field), 318 StopElementHandler (field), 318 xml.parsers.expat.ExpatError (error), 323 xrange (type), 52

yield statement defined, 79 generator expressions, 85

Z

ZeroDivisionError (error), 20, 137

About the Author

Mitchell L Model has worked in a wide range of platforms, languages, technologies, and domains. His specialties include object technology, knowledge representation, user interfaces, distributed computing, and software development practices, and he is an inveterate software tool builder. He has held academic appointments at Brown, Brandeis, and Wesleyan Universities and has worked at a number of early-phase startup companies. As this book went into production, he joined a new synthetic biology startup. For much of his career, he has been an independent consultant, providing training, mentoring, tools, and support to software development groups learning to use new technologies and practices. He has written and taught many professional programming and technology courses and wrote one of the first C++-based data structure books.

Since 1994 Mitchell has been working primarily in bioinformatics, becoming captivated by the complexities of the biological phenomena that it addresses. He was the senior technologist in Millennium Pharmaceutical's large bioinformatics department from 1994 through 2001. In recent years, he has been teaching courses in Northeastern University's Professional Masters in Bioinformatics program.

Mitchell is devoted to teaching, always seeking new ways to effectively communicate the conceptual beauty of computer science. What gives him the greatest satisfaction is coaching students and professionals in the art of software development to enable them to work more effectively and creatively.

Colophon

The animal on the cover of *Bioinformatics Programming Using Python* is a brown rat (*Rattus norvegicus*), also known as a common rat, sewer rat, Norway rat, or wharf rat. One of the largest members of the Muroidea family, the brown rat is 10 to 15 inches long with a 6- to 8-inch long tail. Its fur is coarse and mostly brown or gray. This rat has very sharp hearing and smell, but poor vision. It is omnivorous and nocturnal, and lives almost everywhere humans live, particularly in cities. Like other rodents, brown rats may carry pathogens and spread disease.

Contrary to the brown rat's genus name—norvegicus—it did not originate in Norway. It was named by an 18th-century British naturalist who mistakenly believed that the rat had migrated to England on Norwegian ships in the early 1700s. By the end of the 19th century, scientists had established that the brown rat had most likely originated in China. Today the brown rat has spread to all continents and is the dominant rat in Europe and North America—making it the most successful mammal on earth after humans.

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