Concepts of Programming Languages

ELEVENTH EDITION

Robert W. Sebesta
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Changes for the Eleventh Edition of Concepts of Programming Languages

- **Chapter 6**: Deleted the discussions of Ada’s subrange types, array initialization, records, union types, pointers, and strong typing
- **Chapter 7**: Deleted the discussions of Ada operator associativity and mixed-mode expressions
- **Chapter 8**: Expanded the paragraph on F# selection statements in Section 8.2.1.5
  Deleted the discussion of the Ada `for` statement
- **Chapter 9**: Added three design issues for subprograms in Section 9.3
  Deleted the discussions of Ada and Fortran multidimensional parameters
- **Chapter 10**: Replaced example program `Main_2`, written in Ada, with an equivalent program written in JavaScript in Section 10.4.2
  Changed Figure 10.9 to reflect this new JavaScript example
- **Chapter 11**: Deleted the discussions of Ada abstract data types, generic procedures, and packages
  Added a new paragraph to Section 11.4.3 (Abstract Data Types in Java)
- **Chapter 12**: Added a paragraph to Section 12.2.2 (Inheritance) that discusses access control
  Expanded the discussion of class variables in Section 12.2.2
  Added a paragraph to Section 12.4.4 that discusses final classes in Objective-C
  Reorganized Sections 12.5 to 12.9 into a single section
  Added Table 12.1 on language design choices to Section 12.4.6.4
  Added a new section, Section 6 (Reflection), including example programs in Java and C#
- **Chapter 13**: Deleted the discussions of Ada task termination and task priorities
- **Chapter 14**: Deleted exception handling in Ada
  Added a new section, 14.4 (Exception Handling in Python and Ruby)
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Preface

Changes for the Eleventh Edition

The goals, overall structure, and approach of this eleventh edition of *Concepts of Programming Languages* remain the same as those of the ten earlier editions. The principal goals are to introduce the fundamental constructs of contemporary programming languages and to provide the reader with the tools necessary for the critical evaluation of existing and future programming languages. A secondary goal is to prepare the reader for the study of compiler design, by providing an in-depth discussion of programming language structures, presenting a formal method of describing syntax, and introducing approaches to lexical and syntactic analysis.

The eleventh edition evolved from the tenth through several different kinds of changes. To maintain the currency of the material, much of the discussion of older programming languages, particularly Ada and Fortran, has been removed. For example, the descriptions of Ada’s records, union types, and pointers were removed from Chapter 6. Likewise, the description of Ada’s *for* statement was removed from Chapter 8 and the discussion of Ada’s abstract data types was removed from Chapter 11.

On the other hand, a section on reflection that includes two complete program examples was added to Chapter 12, a section on exception handling in Python and Ruby was added to Chapter 14, and a table of the design choices of a few common languages for support for object-oriented programming was added to Chapter 12.

In some cases, material has been moved. For example, Section 9.10 was moved backward to become the new Section 9.8.

In one case, example program *MAIN_2* in Chapter 10 was rewritten in JavaScript, previously having been written in Ada.

Chapter 12 was substantially revised, with several new paragraphs, two new sections, and numerous other changes to improve clarity.

The Vision

This book describes the fundamental concepts of programming languages by discussing the design issues of the various language constructs, examining the design choices for these constructs in some of the most common languages, and critically comparing design alternatives.
Any serious study of programming languages requires an examination of some related topics, among which are formal methods of describing the syntax and semantics of programming languages, which are covered in Chapter 3. Also, implementation techniques for various language constructs must be considered: Lexical and syntax analysis are discussed in Chapter 4, and implementation of subprogram linkage is covered in Chapter 10. Implementation of some other language constructs is discussed in various other parts of the book.

The following paragraphs outline the contents of the eleventh edition.

Chapter Outlines

Chapter 1 begins with a rationale for studying programming languages. It then discusses the criteria used for evaluating programming languages and language constructs. The primary influences on language design, common design trade-offs, and the basic approaches to implementation are also examined.

Chapter 2 outlines the evolution of the languages that are discussed in this book. Although no attempt is made to describe any language completely, the origins, purposes, and contributions of each are discussed. This historical overview is valuable, because it provides the background necessary to understanding the practical and theoretical basis for contemporary language design. It also motivates further study of language design and evaluation. Because none of the remainder of the book depends on Chapter 2, it can be read on its own, independent of the other chapters.

Chapter 3 describes the primary formal method for describing the syntax of programming language—BNF. This is followed by a description of attribute grammars, which describe both the syntax and static semantics of languages. The difficult task of semantic description is then explored, including brief introductions to the three most common methods: operational, denotational, and axiomatic semantics.

Chapter 4 introduces lexical and syntax analysis. This chapter is targeted to those Computer Science departments that no longer require a compiler design course in their curricula. Similar to Chapter 2, this chapter stands alone and can be studied independently of the rest of the book, except for Chapter 3, on which it depends.

Chapters 5 through 14 describe in detail the design issues for the primary constructs of programming languages. In each case, the design choices for several example languages are presented and evaluated. Specifically, Chapter 5 covers the many characteristics of variables, Chapter 6 covers data types, and Chapter 7 explains expressions and assignment statements. Chapter 8 describes control statements, and Chapters 9 and 10 discuss subprograms and their implementation. Chapter 11 examines data abstraction facilities. Chapter 12 provides an in-depth discussion of language features that support object-oriented programming (inheritance and dynamic method binding), Chapter 13 discusses concurrent program units, and Chapter 14 is about exception handling, along with a brief discussion of event handling.
Chapters 15 and 16 describe two of the most important alternative programming paradigms: functional programming and logic programming. However, some of the data structures and control constructs of functional programming languages are discussed in Chapters 6 and 8. Chapter 15 presents an introduction to Scheme, including descriptions of some of its primitive functions, special forms, and functional forms, as well as some examples of simple functions written in Scheme. Brief introductions to ML, Haskell, and F# are given to illustrate some different directions in functional language design. Chapter 16 introduces logic programming and the logic programming language, Prolog.

To the Instructor

In the junior-level programming language course at the University of Colorado at Colorado Springs, the book is used as follows: We typically cover Chapters 1 and 3 in detail, and though students find it interesting and beneficial reading, Chapter 2 receives little lecture time due to its lack of hard technical content. Because no material in subsequent chapters depends on Chapter 2, as noted earlier, it can be skipped entirely, and because we require a course in compiler design, Chapter 4 is not covered.

Chapters 5 through 9 should be relatively easy for students with extensive programming experience in C++, Java, or C#. Chapters 10 through 14 are more challenging and require more detailed lectures.

Chapters 15 and 16 are entirely new to most students at the junior level. Ideally, language processors for Scheme and Prolog should be available for students required to learn the material in these chapters. Sufficient material is included to allow students to dabble with some simple programs.

Undergraduate courses will probably not be able to cover all of the material in the last two chapters. Graduate courses, however, should be able to completely discuss the material in those chapters by skipping over some parts of the early chapters on imperative languages.

Supplemental Materials

The following supplements are available to all readers of this book at www.pearsonglobaleditions.com/Sebesta.

- A set of lecture note slides. PowerPoint slides are available for each chapter in the book.
- All of the figures from the book.

A companion Web site to the book is available at www.pearsonglobaleditions.com/Sebesta. This site contains mini-manuals (approximately 100-page tutorials) on a handful of languages. These assume that the student knows how to program
in some other language, giving the student enough information to complete the chapter materials in each language. Currently the site includes manuals for C++, C, Java, and Smalltalk.

Solutions to many of the problem sets are available to qualified instructors in our Instructor Resource Center at www.pearsonglobaleditions.com/Sebesta.

**Language Processor Availability**

Processors for and information about some of the programming languages discussed in this book can be found at the following Web sites:

- C, C++, Fortran, and Ada: gcc.gnu.org
- C# and F#: microsoft.com
- Java: java.sun.com
- Haskell: haskell.org
- Lua: www.lua.org
- Scheme: www.plt-scheme.org/software/drscheme
- Perl: www.perl.com
- Python: www.python.org
- Ruby: www.ruby-lang.org

JavaScript is included in virtually all browsers; PHP is included in virtually all Web servers.

All this information is also included on the companion Web site.
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CONCEPTS OF
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Preliminaries

1.1 Reasons for Studying Concepts of Programming Languages
1.2 Programming Domains
1.3 Language Evaluation Criteria
1.4 Influences on Language Design
1.5 Language Categories
1.6 Language Design Trade-Offs
1.7 Implementation Methods
1.8 Programming Environments
Before we begin discussing the concepts of programming languages, we must consider a few preliminaries. First, we explain some reasons why computer science students and professional software developers should study general concepts of language design and evaluation. This discussion is especially valuable for those who believe that a working knowledge of one or two programming languages is sufficient for computer scientists. Then, we briefly describe the major programming domains. Next, because the book evaluates language constructs and features, we present a list of criteria that can serve as a basis for such judgments. Then, we discuss the two major influences on language design: machine architecture and program design methodologies. After that, we introduce the various categories of programming languages. Next, we describe a few of the major trade-offs that must be considered during language design.

Because this book is also about the implementation of programming languages, this chapter includes an overview of the most common general approaches to implementation. Finally, we briefly describe a few examples of programming environments and discuss their impact on software production.

1.1 Reasons for Studying Concepts of Programming Languages

It is natural for students to wonder how they will benefit from the study of programming language concepts. After all, many other topics in computer science are worthy of serious study. The following is what we believe to be a compelling list of potential benefits of studying concepts of programming languages:

- **Increased capacity to express ideas.** It is widely believed that the depth at which people can think is influenced by the expressive power of the language in which they communicate their thoughts. Those with only a weak understanding of natural language are limited in the complexity of their thoughts, particularly in depth of abstraction. In other words, it is difficult for people to conceptualize structures they cannot describe, verbally or in writing.

  Programmers, in the process of developing software, are similarly constrained. The language in which they develop software places limits on the kinds of control structures, data structures, and abstractions they can use; thus, the forms of algorithms they can construct are likewise limited. Awareness of a wider variety of programming language features can reduce such limitations in software development. Programmers can increase the range of their software development thought processes by learning new language constructs.

  It might be argued that learning the capabilities of other languages does not help a programmer who is forced to use a language that lacks those capabilities. That argument does not hold up, however, because often, language constructs can be simulated in other languages that do not support those constructs directly. For example, a C programmer who had learned the structure and uses of associative arrays in Perl (Christianson et al., 2012) might design structures that simulate associative arrays in that language. In other words, the study of programming language concepts builds an
appreciation for valuable language features and constructs and encourages programmers to use them, even when the language they are using does not directly support such features and constructs.

- **Improved background for choosing appropriate languages.** Some professional programmers have had little formal education in computer science; rather, they have developed their programming skills independently or through in-house training programs. Such training programs often limit instruction to one or two languages that are directly relevant to the current projects of the organization. Other programmers received their formal training years ago. The languages they learned then are no longer used, and many features now available in programming languages were not widely known at the time. The result is that many programmers, when given a choice of languages for a new project, use the language with which they are most familiar, even if it is poorly suited for the project at hand. If these programmers were familiar with a wider range of languages and language constructs, they would be better able to choose the language with the features that best address the problem.

Some of the features of one language often can be simulated in another language. However, it is preferable to use a feature whose design has been integrated into a language than to use a simulation of that feature, which is often less elegant, more cumbersome, and less safe.

- **Increased ability to learn new languages.** Computer programming is still a relatively young discipline, and design methodologies, software development tools, and programming languages are still in a state of continuous evolution. This makes software development an exciting profession, but it also means that continuous learning is essential. The process of learning a new programming language can be lengthy and difficult, especially for someone who is comfortable with only one or two languages and has never examined programming language concepts in general. Once a thorough understanding of the fundamental concepts of languages is acquired, it becomes far easier to see how these concepts are incorporated into the design of the language being learned. For example, programmers who understand the concepts of object-oriented programming will have a much easier time learning Ruby (Thomas et al., 2013) than those who have never used those concepts.

The same phenomenon occurs in natural languages. The better you know the grammar of your native language, the easier it is to learn a second language. Furthermore, learning a second language has the benefit of teaching you more about your first language.

The TIOBE Programming Community issues an index (http://www.tiobe.com/index.php/content/paperinfo/tpci/index.htm) that is an indicator of the relative popularity of programming languages. For example, according to the index, C, Java, and Objective-C were the three most popular languages in use in February 2014.\(^1\) However, dozens of other

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\(^1\) Note that this index is only one measure of the popularity of programming languages, and its accuracy is not universally accepted.
languages were widely used at the time. The index data also show that the distribution of usage of programming languages is always changing. The number of languages in use and the dynamic nature of the statistics imply that every software developer must be prepared to learn different languages.

Finally, it is essential that practicing programmers know the vocabulary and fundamental concepts of programming languages so they can read and understand programming language descriptions and evaluations, as well as promotional literature for languages and compilers. These are the sources of information needed in order to choose and learn a language.

- **Better understanding of the significance of implementation.** In learning the concepts of programming languages, it is both interesting and necessary to touch on the implementation issues that affect those concepts. In some cases, an understanding of implementation issues leads to an understanding of why languages are designed the way they are. In turn, this knowledge leads to the ability to use a language more intelligently, as it was designed to be used. We can become better programmers by understanding the choices among programming language constructs and the consequences of those choices.

  Certain kinds of program bugs can be found and fixed only by a programmer who knows some related implementation details. Another benefit of understanding implementation issues is that it allows us to visualize how a computer executes various language constructs. In some cases, some knowledge of implementation issues provides hints about the relative efficiency of alternative constructs that may be chosen for a program. For example, programmers who know little about the complexity of the implementation of subprogram calls often do not realize that a small subprogram that is frequently called can be a highly inefficient design choice.

  Because this book touches on only a few of the issues of implementation, the previous two paragraphs also serve well as rationale for studying compiler design.

- **Better use of languages that are already known.** Most contemporary programming languages are large and complex. Accordingly, it is uncommon for a programmer to be familiar with and use all of the features of a language he or she uses. By studying the concepts of programming languages, programmers can learn about previously unknown and unused parts of the languages they already use and begin to use those features.

- **Overall advancement of computing.** Finally, there is a global view of computing that can justify the study of programming language concepts. Although it is usually possible to determine why a particular programming language became popular, many believe, at least in retrospect, that the most popular languages are not always the best available. In some cases, it might be concluded that a language became widely used, at least in part, because those in positions to choose languages were not sufficiently familiar with programming language concepts.

  For example, many people believe it would have been better if ALGOL 60 (Backus et al., 1963) had displaced Fortran (McCracken, 1961) in the
1.2 Programming Domains

Computers have been applied to a myriad of different areas, from controlling nuclear power plants to providing video games in mobile phones. Because of this great diversity in computer use, programming languages with very different goals have been developed. In this section, we briefly discuss a few of the areas of computer applications and their associated languages.

1.2.1 Scientific Applications

The first digital computers, which appeared in the late 1940s and early 1950s, were invented and used for scientific applications. Typically, the scientific applications of that time used relatively simple data structures, but required large numbers of floating-point arithmetic computations. The most common data structures were arrays and matrices; the most common control structures were counting loops and selections. The early high-level programming languages invented for scientific applications were designed to provide for those needs. Their competition was assembly language, so efficiency was a primary concern. The first language for scientific applications was Fortran. ALGOL 60 and most of its descendants were also intended to be used in this area, although they were designed to be used in related areas as well. For some scientific applications where efficiency is the primary concern, such as those that were common in the 1950s and 1960s, no subsequent language is significantly better than Fortran, which explains why Fortran is still used.

1.2.2 Business Applications

The use of computers for business applications began in the 1950s. Special computers were developed for this purpose, along with special languages. The first successful high-level language for business was COBOL (ISO/IEC, 2002),
the initial version of which appeared in 1960. It probably still is the most commonly used language for these applications. Business languages are characterized by facilities for producing elaborate reports, precise ways of describing and storing decimal numbers and character data, and the ability to specify decimal arithmetic operations.

There have been few developments in business application languages outside the development and evolution of COBOL. Therefore, this book includes only limited discussions of the structures in COBOL.

### 1.2.3 Artificial Intelligence

Artificial intelligence (AI) is a broad area of computer applications characterized by the use of symbolic rather than numeric computations. Symbolic computation means that symbols, consisting of names rather than numbers, are manipulated. Also, symbolic computation is more conveniently done with linked lists of data rather than arrays. This kind of programming sometimes requires more flexibility than other programming domains. For example, in some AI applications the ability to create and execute code segments during execution is convenient.

The first widely used programming language developed for AI applications was the functional language Lisp (McCarthy et al., 1965), which appeared in 1959. Most AI applications developed prior to 1990 were written in Lisp or one of its close relatives. During the early 1970s, however, an alternative approach to some of these applications appeared—logic programming using the Prolog (Clocksin and Mellish, 2013) language. More recently, some AI applications have been written in systems languages such as C. Scheme (Dybvig, 2009), a dialect of Lisp, and Prolog are introduced in Chapters 15 and 16, respectively.

### 1.2.4 Web Software

The World Wide Web is supported by an eclectic collection of languages, ranging from markup languages, such as HTML, which is not a programming language, to general-purpose programming languages, such as Java. Because of the pervasive need for dynamic Web content, some computation capability is often included in the technology of content presentation. This functionality can be provided by embedding programming code in an HTML document. Such code is often in the form of a scripting language, such as JavaScript or PHP (Tatroe, 2013). There are also some markup-like languages that have been extended to include constructs that control document processing, which are discussed in Section 1.5 and in Chapter 2.

### 1.3 Language Evaluation Criteria

As noted previously, the purpose of this book is to examine carefully the underlying concepts of the various constructs and capabilities of programming languages. We will also evaluate these features, focusing on their impact on the
software development process, including maintenance. To do this, we need a set of evaluation criteria. Such a list of criteria is necessarily controversial, because it is difficult to get even two computer scientists to agree on the value of some given language characteristic relative to others. In spite of these differences, most would agree that the criteria discussed in the following subsections are important.

Some of the characteristics that influence three of the four most important of these criteria are shown in Table 1.1, and the criteria themselves are discussed in the following sections. Note that only the most important characteristics are included in the table, mirroring the discussion in the following subsections. One could probably make the case that if one considered less important characteristics, virtually all table positions could include “bullets.”

Note that some of these characteristics are broad and somewhat vague, such as writability, whereas others are specific language constructs, such as exception handling. Furthermore, although the discussion might seem to imply that the criteria have equal importance, that implication is not intended, and it is clearly not the case.

### 1.3.1 Readability

One of the most important criteria for judging a programming language is the ease with which programs can be read and understood. Before 1970, software development was largely thought of in terms of writing code. The primary positive characteristic of programming languages was efficiency. Language constructs were designed more from the point of view of the computer than of the computer users. In the 1970s, however, the software life-cycle concept (Booch, 1987) was developed; coding was relegated to a much smaller role,
and maintenance was recognized as a major part of the cycle, particularly in terms of cost. Because ease of maintenance is determined in large part by the readability of programs, readability became an important measure of the quality of programs and programming languages. This was an important juncture in the evolution of programming languages. There was a distinct crossover from a focus on machine orientation to a focus on human orientation.

Readability must be considered in the context of the problem domain. For example, if a program that describes a computation is written in a language not designed for such use, the program may be unnatural and convoluted, making it unusually difficult to read.

The following subsections describe characteristics that contribute to the readability of a programming language.

1.3.1.1 Overall Simplicity

The overall simplicity of a programming language strongly affects its readability. A language with a large number of basic constructs is more difficult to learn than one with a smaller number. Programmers who must use a large language often learn a subset of the language and ignore its other features. This learning pattern is sometimes used to excuse the large number of language constructs, but that argument is not valid. Readability problems occur whenever the program’s author has learned a different subset from that subset with which the reader is familiar.

A second complicating characteristic of a programming language is feature multiplicity—that is, having more than one way to accomplish a particular operation. For example, in Java, a user can increment a simple integer variable in four different ways:

\[
\begin{align*}
\text{count} & = \text{count} + 1 \\
\text{count} & += 1 \\
\text{count} & ++ \\
++\text{count}
\end{align*}
\]

Although the last two statements have slightly different meanings from each other and from the others in some contexts, all of them have the same meaning when used as stand-alone expressions. These variations are discussed in Chapter 7.

A third potential problem is operator overloading, in which a single operator symbol has more than one meaning. Although this is often useful, it can lead to reduced readability if users are allowed to create their own overloading and do not do it sensibly. For example, it is clearly acceptable to overload + to use it for both integer and floating-point addition. In fact, this overloading simplifies a language by reducing the number of operators. However, suppose the programmer defined + used between single-dimensioned array operands to mean the sum of all elements of both arrays. Because the usual meaning of vector addition is quite different from this, this unusual meaning could confuse
both the author and the program’s readers. An even more extreme example of program confusion would be a user defining + between two vector operands to mean the difference between their respective first elements. Operator overloading is further discussed in Chapter 7.

Simplicity in languages can, of course, be carried too far. For example, the form and meaning of most assembly language statements are models of simplicity, as you can see when you consider the statements that appear in the next section. This very simplicity, however, makes assembly language programs less readable. Because they lack more complex control statements, program structure is less obvious; because the statements are simple, far more of them are required than in equivalent programs in a high-level language. These same arguments apply to the less extreme case of high-level languages with inadequate control and data-structuring constructs.

1.3.1.2 Orthogonality

Orthogonality in a programming language means that a relatively small set of primitive constructs can be combined in a relatively small number of ways to build the control and data structures of the language. Furthermore, every possible combination of primitives is legal and meaningful. For example, consider data types. Suppose a language has four primitive data types (integer, float, double, and character) and two type operators (array and pointer). If the two type operators can be applied to themselves and the four primitive data types, a large number of data structures can be defined.

The meaning of an orthogonal language feature is independent of the context of its appearance in a program. (The word orthogonal comes from the mathematical concept of orthogonal vectors, which are independent of each other.) Orthogonality follows from a symmetry of relationships among primitives. A lack of orthogonality leads to exceptions to the rules of the language. For example, in a programming language that supports pointers, it should be possible to define a pointer to point to any specific type defined in the language. However, if pointers are not allowed to point to arrays, many potentially useful user-defined data structures cannot be defined.

We can illustrate the use of orthogonality as a design concept by comparing one aspect of the assembly languages of the IBM mainframe computers and the VAX series of minicomputers. We consider a single simple situation: adding two 32-bit integer values that reside in either memory or registers and replacing one of the two values with the sum. The IBM mainframes have two instructions for this purpose, which have the forms

\[
\begin{align*}
A & \text{ Reg1, memory}_\text{cell} \\
\text{AR} & \text{ Reg1, Reg2}
\end{align*}
\]

where Reg1 and Reg2 represent registers. The semantics of these are

\[
\begin{align*}
\text{Reg1} & \leftarrow \text{contents(Reg1)} + \text{contents(memory}_\text{cell}) \\
\text{Reg1} & \leftarrow \text{contents(Reg1)} + \text{contents(Reg2)}
\end{align*}
\]
The VAX addition instruction for 32-bit integer values is

\texttt{ADDL \text{operand\_1}, \text{operand\_2}}

whose semantics is

\texttt{\text{operand\_2} \leftarrow \text{contents}(\text{operand\_1}) + \text{contents}(\text{operand\_2})}

In this case, either operand can be a register or a memory cell.

The VAX instruction design is orthogonal in that a single instruction can use either registers or memory cells as the operands. There are two ways to specify operands, which can be combined in all possible ways. The IBM design is not orthogonal. Only two out of four operand combinations possibilities are legal, and the two require different instructions, \texttt{A} and \texttt{AR}. The IBM design is more restricted and therefore less writable. For example, you cannot add two values and store the sum in a memory location. Furthermore, the IBM design is more difficult to learn because of the restrictions and the additional instruction.

Orthogonality is closely related to simplicity: The more orthogonal the design of a language, the fewer exceptions the language rules require. Fewer exceptions mean a higher degree of regularity in the design, which makes the language easier to learn, read, and understand. Anyone who has learned a significant part of the English language can testify to the difficulty of learning its many rule exceptions (for example, \textit{i} before \textit{e} except after \textit{c}).

As examples of the lack of orthogonality in a high-level language, consider the following rules and exceptions in C. Although C has two kinds of structured data types, arrays and records (\texttt{struct}s), records can be returned from functions but arrays cannot. A member of a structure can be any data type except \texttt{void} or a structure of the same type. An array element can be any data type except \texttt{void} or a function. Parameters are passed by value, unless they are arrays, in which case they are, in effect, passed by reference (because the appearance of an array name without a subscript in a C program is interpreted to be the address of the array’s first element).

As an example of context dependence, consider the C expression

\texttt{a + b}

This expression often means that the values of \texttt{a} and \texttt{b} are fetched and added together. However, if \texttt{a} happens to be a pointer and \texttt{b} is an integer, it affects the value of \texttt{b}. For example, if \texttt{a} points to a float value that occupies four bytes, then the value of \texttt{b} must be scaled—in this case multiplied by 4—before it is added to \texttt{a}. Therefore, the type of \texttt{a} affects the treatment of the value of \texttt{b}. The context of \texttt{b} affects its meaning.

Too much orthogonality can also cause problems. Perhaps the most orthogonal programming language is ALGOL 68 (van Wijngaarden et al., 1969). Every language construct in ALGOL 68 has a type, and there are no restrictions on those types. In addition, most constructs produce values. This combinational freedom allows extremely complex constructs. For example, a
conditional can appear as the left side of an assignment, along with declarations
and other assorted statements, as long as the result is an address. This extreme
form of orthogonality leads to unnecessary complexity. Furthermore, because
languages require a large number of primitives, a high degree of orthogonality
results in an explosion of combinations. So, even if the combinations are simple,
their sheer numbers lead to complexity.

Simplicity in a language, therefore, is at least in part the result of a combi-
nation of a relatively small number of primitive constructs and a limited use of
the concept of orthogonality.

Some believe that functional languages offer a good combination of simplic-
ity and orthogonality. A functional language, such as Lisp, is one in which comput-
ations are made primarily by applying functions to given parameters. In contrast,
in imperative languages such as C, C++, and Java, computations are usually speci-
ﬁed with variables and assignment statements. Functional languages offer poten-
tially the greatest overall simplicity, because they can accomplish everything with
a single construct, the function call, which can be combined simply with other
function calls. This simple elegance is the reason why some language researchers
are attracted to functional languages as the primary alternative to complex non-
functional languages such as Java. Other factors, such as efﬁciency, however, have
prevented functional languages from becoming more widely used.

1.3.1.3 Data Types

The presence of adequate facilities for deﬁning data types and data structures
in a language is another signiﬁcant aid to readability. For example, suppose a
numeric type is used for an indicator ﬂag because there is no Boolean type in
the language. In such a language, we might have an assignment such as the
following:

timeOut = 1

The meaning of this statement is unclear, whereas in a language that includes
Boolean types, we would have the following:

timeOut = true

The meaning of this statement is perfectly clear.

1.3.1.4 Syntax Design

The syntax, or form, of the elements of a language has a signiﬁcant effect on
the readability of programs. Following are some examples of syntactic design
choices that affect readability:

• Special words. Program appearance and thus program readability are
  strongly inﬂuenced by the forms of a language’s special words (for example,
  while, class, and for). Especially important is the method of forming
compound statements, or statement groups, primarily in control constructs. Some languages have used matching pairs of special words or symbols to form groups. C and its descendants use braces to specify compound statements. All of these languages have diminished readability because statement groups are always terminated in the same way, which makes it difficult to determine which group is being ended when an end or a right brace appears. Fortran 95 and Ada (ISO/IEC, 2014) make this clearer by using a distinct closing syntax for each type of statement group. For example, Ada uses end if to terminate a selection construct and end loop to terminate a loop construct. This is an example of the conflict between simplicity that results in fewer reserved words, as in Java, and the greater readability that can result from using more reserved words, as in Ada.

Another important issue is whether the special words of a language can be used as names for program variables. If so, the resulting programs can be very confusing. For example, in Fortran 95, special words, such as Do and End, are legal variable names, so the appearance of these words in a program may or may not connote something special.

- **Form and meaning.** Designing statements so that their appearance at least partially indicates their purpose is an obvious aid to readability. Semantics, or meaning, should follow directly from syntax, or form. In some cases, this principle is violated by two language constructs that are identical or similar in appearance but have different meanings, depending perhaps on context. In C, for example, the meaning of the reserved word static depends on the context of its appearance. If used on the definition of a variable inside a function, it means the variable is created at compile time. If used on the definition of a variable that is outside all functions, it means the variable is visible only in the file in which its definition appears; that is, it is not exported from that file.

One of the primary complaints about the shell commands of UNIX (Robbins, 2005) is that their appearance does not always suggest their function. For example, the meaning of the UNIX command grep can be deciphered only through prior knowledge, or perhaps cleverness and familiarity with the UNIX editor, ed. The appearance of grep connotes nothing to UNIX beginners. (In ed, the command /regular_expression/ searches for a substring that matches the regular expression. Preceding this with g makes it a global command, specifying that the scope of the search is the whole file being edited. Following the command with p specifies that lines with the matching substring are to be printed. So g/regular_expression/p, which can obviously be abbreviated as grep, prints all lines in a file that contain substrings that match the regular expression.)

### 1.3.2 Writability

Writability is a measure of how easily a language can be used to create programs for a chosen problem domain. Most of the language characteristics that affect readability also affect writability. This follows directly from the fact that the
process of writing a program requires the programmer frequently to reread the part of the program that is already written.

As is the case with readability, writability must be considered in the context of the target problem domain of a language. It simply is not fair to compare the writability of two languages in the realm of a particular application when one was designed for that application and the other was not. For example, the writabilities of Visual BASIC (VB) (Halvorson, 2013) and C are dramatically different for creating a program that has a graphical user interface (GUI), for which VB is ideal. Their writabilities are also quite different for writing systems programs, such as an operation system, for which C was designed.

The following subsections describe the most important characteristics influencing the writability of a language.

1.3.2.1 Simplicity and Orthogonality

If a language has a large number of different constructs, some programmers might not be familiar with all of them. This situation can lead to a misuse of some features and a disuse of others that may be either more elegant or more efficient, or both, than those that are used. It may even be possible, as noted by Hoare (1973), to use unknown features accidentally, with bizarre results. Therefore, a smaller number of primitive constructs and a consistent set of rules for combining them (that is, orthogonality) is much better than simply having a large number of primitives. A programmer can design a solution to a complex problem after learning only a simple set of primitive constructs.

On the other hand, too much orthogonality can be a detriment to writability. Errors in programs can go undetected when nearly any combination of primitives is legal. This can lead to code absurdities that cannot be discovered by the compiler.

1.3.2.2 Expressivity

Expressivity in a language can refer to several different characteristics. In a language such as APL (Gilman and Rose, 1983), it means that there are very powerful operators that allow a great deal of computation to be accomplished with a very small program. More commonly, it means that a language has relatively convenient, rather than cumbersome, ways of specifying computations. For example, in C, the notation `count++` is more convenient and shorter than `count = count + 1`. Also, the `and then` Boolean operator in Ada is a convenient way of specifying short-circuit evaluation of a Boolean expression. The inclusion of the `for` statement in Java makes writing counting loops easier than with the use of `while`, which is also possible. All of these increase the writability of a language.

1.3.3 Reliability

A program is said to be reliable if it performs to its specifications under all conditions. The following subsections describe several language features that have a significant effect on the reliability of programs in a given language.
1.3.3.1 Type Checking

Type checking is simply testing for type errors in a given program, either by the compiler or during program execution. Type checking is an important factor in language reliability. Because run-time type checking is expensive, compile-time type checking is more desirable. Furthermore, the earlier errors in programs are detected, the less expensive it is to make the required repairs. The design of Java requires checks of the types of nearly all variables and expressions at compile time. This virtually eliminates type errors at run time in Java programs. Types and type checking are discussed in depth in Chapter 6.

One example of how failure to type check, at either compile time or run time, has led to countless program errors is the use of subprogram parameters in the original C language (Kernighan and Ritchie, 1978). In this language, the type of an actual parameter in a function call was not checked to determine whether its type matched that of the corresponding formal parameter in the function. An int type variable could be used as an actual parameter in a call to a function that expected a float type as its formal parameter, and neither the compiler nor the run-time system would detect the inconsistency. For example, because the bit string that represents the integer 23 is essentially unrelated to the bit string that represents a floating-point 23, if an integer 23 is sent to a function that expects a floating-point parameter, any uses of the parameter in the function will produce nonsense. Furthermore, such problems are often difficult to diagnose. The current version of C has eliminated this problem by requiring all parameters to be type checked. Subprograms and parameter-passing techniques are discussed in Chapter 9.

1.3.3.2 Exception Handling

The ability of a program to intercept run-time errors (as well as other unusual conditions detectable by the program), take corrective measures, and then continue is an obvious aid to reliability. This language facility is called exception handling. Ada, C++, Java, and C# include extensive capabilities for exception handling, but such facilities are practically nonexistent in some widely used languages, for example C. Exception handling is discussed in Chapter 14.

1.3.3.3 Aliasing

Loosely defined, aliasing is having two or more distinct names in a program that can be used to access the same memory cell. It is now generally accepted that aliasing is a dangerous feature in a programming language. Most programming languages allow some kind of aliasing—for example, two pointers (or references) set to point to the same variable, which is possible in most languages. In such a program, the programmer must always remember that changing the

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3. In response to this and other similar problems, UNIX systems include a utility program named lint that checks C programs for such problems.
value pointed to by one of the two changes the value referenced by the other. Some kinds of aliasing, as described in Chapters 5 and 9, can be prohibited by the design of a language.

In some languages, aliasing is used to overcome deficiencies in the language’s data abstraction facilities. Other languages greatly restrict aliasing to increase their reliability.

### 1.3.3.4 Readability and Writability

Both readability and writability influence reliability. A program written in a language that does not support natural ways to express the required algorithms will necessarily use unnatural approaches. Unnatural approaches are less likely to be correct for all possible situations. The easier a program is to write, the more likely it is to be correct.

Readability affects reliability in both the writing and maintenance phases of the life cycle. Programs that are difficult to read are difficult both to write and to modify.

### 1.3.4 Cost

The total cost of a programming language is a function of many of its characteristics.

First, there is the cost of training programmers to use the language, which is a function of the simplicity and orthogonality of the language and the experience of the programmers. Although more powerful languages are not necessarily more difficult to learn, they often are.

Second, there is the cost of writing programs in the language. This is a function of the writability of the language, which depends in part on its closeness in purpose to the particular application. The original efforts to design and implement high-level languages were driven by the desire to lower the costs of creating software.

Both the cost of training programmers and the cost of writing programs in a language can be significantly reduced in a good programming environment. Programming environments are discussed in Section 1.8.

Third, there is the cost of compiling programs in the language. A major impediment to the early use of Ada was the prohibitively high cost of running the first-generation Ada compilers. This problem was diminished by the appearance of improved Ada compilers.

Fourth, the cost of executing programs written in a language is greatly influenced by that language’s design. A language that requires many run-time type checks will prohibit fast code execution, regardless of the quality of the compiler. Although execution efficiency was the foremost concern in the design of early languages, it is now considered to be less important.

A simple trade-off can be made between compilation cost and execution speed of the compiled code. **Optimization** is the name given to the collection of techniques that compilers may use to decrease the size and/or increase the
execution speed of the code they produce. If little or no optimization is done, compilation can be done much faster than if a significant effort is made to produce optimized code. The choice between the two alternatives is influenced by the environment in which the compiler will be used. In a laboratory for beginning programming students, who often compile their programs many times during development but use little code at execution time (their programs are small and they must execute correctly only once), little or no optimization should be done. In a production environment, where compiled programs are executed many times after development, it is better to pay the extra cost to optimize the code.

The fifth factor in the cost of a language is the cost of the language implementation system. One of the factors that explains the rapid acceptance of Java is that free compiler/interpreter systems became available for it soon after its design was released. A language whose implementation system is either expensive or runs only on expensive hardware will have a much smaller chance of becoming widely used. For example, the high cost of first-generation Ada compilers helped prevent Ada from becoming popular in its early days.

Sixth, there is the cost of poor reliability. If the software fails in a critical system, such as a nuclear power plant or an X-ray machine for medical use, the cost could be very high. The failures of noncritical systems can also be very expensive in terms of lost future business or lawsuits over defective software systems.

The final consideration is the cost of maintaining programs, which includes both corrections and modifications to add new functionality. The cost of software maintenance depends on a number of language characteristics, primarily readability. Because maintenance is often done by individuals other than the original author of the software, poor readability can make the task extremely challenging.

The importance of software maintainability cannot be overstated. It has been estimated that for large software systems with relatively long lifetimes, maintenance costs can be as high as two to four times as much as development costs (Sommerville, 2010).

Of all the contributors to language costs, three are most important: program development, maintenance, and reliability. Because these are functions of writability and readability, these two evaluation criteria are, in turn, the most important.

Of course, a number of other criteria could be used for evaluating programming languages. One example is portability, or the ease with which programs can be moved from one implementation to another. Portability is most strongly influenced by the degree of standardization of the language. Some languages are not standardized at all, making programs in these languages very difficult to move from one implementation to another. This problem is alleviated in some cases by the fact that implementations for some languages now have single sources. Standardization is a time-consuming and difficult process. A committee began work on producing a standard version of C++ in 1989. It was approved in 1998.
Generality (the applicability to a wide range of applications) and well-definedness (the completeness and precision of the language’s official defining document) are two other criteria.

Most criteria, particularly readability, writability, and reliability, are neither precisely defined nor precisely measurable. Nevertheless, they are useful concepts and they provide valuable insight into the design and evaluation of programming languages.

A final note on evaluation criteria: language design criteria are weighed differently from different perspectives. Language implementors are concerned primarily with the difficulty of implementing the constructs and features of the language. Language users are worried about writability first and readability later. Language designers are likely to emphasize elegance and the ability to attract widespread use. These characteristics often conflict with one another.

### 1.4 Influences on Language Design

In addition to those factors described in Section 1.3, several other factors influence the basic design of programming languages. The most important of these are computer architecture and programming design methodologies.

#### 1.4.1 Computer Architecture

The basic architecture of computers has had a profound effect on language design. Most of the popular languages of the past 60 years have been designed around the prevalent computer architecture, called the von Neumann architecture, after one of its originators, John von Neumann (pronounced “von Noyman”). These languages are called imperative languages. In a von Neumann computer, both data and programs are stored in the same memory. The central processing unit (CPU), which executes instructions, is separate from the memory. Therefore, instructions and data must be transmitted, or piped, from memory to the CPU. Results of operations in the CPU must be moved back to memory. Nearly all digital computers built since the 1940s have been based on the von Neumann architecture. The overall structure of a von Neumann computer is shown in Figure 1.1.

Because of the von Neumann architecture, the central features of imperative languages are variables, which model the memory cells; assignment statements, which are based on the piping operation; and the iterative form of repetition, which is the most efficient way to implement repetition on this architecture. Operands in expressions are piped from memory to the CPU, and the result of evaluating the expression is piped back to the memory cell represented by the left side of the assignment. Iteration is fast on von Neumann computers because instructions are stored in adjacent cells of memory and repeating the execution of a section of code requires only a branch instruction. This efficiency discourages the use of recursion for repetition, although recursion is sometimes more natural.
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The execution of a machine code program on a von Neumann architecture computer occurs in a process called the *fetch-execute cycle*. As stated earlier, programs reside in memory but are executed in the CPU. Each instruction to be executed must be moved from memory to the processor. The address of the next instruction to be executed is maintained in a register called the *program counter*. The fetch-execute cycle can be simply described by the following algorithm:

initialize the program counter
repeat forever
    fetch the instruction pointed to by the program counter
    increment the program counter to point at the next instruction
    decode the instruction
    execute the instruction
end repeat

The “decode the instruction” step in the algorithm means the instruction is examined to determine what action it specifies. Program execution terminates when a stop instruction is encountered, although on an actual computer a stop instruction is rarely executed. Rather, control transfers from the operating system to a user program for its execution and then back to the operating system when the user program execution is complete. In a computer system in which more than one user program may be in memory at a given time, this process is far more complex.

As stated earlier, a functional, or applicative, language is one in which the primary means of computation is applying functions to given parameters. Programming can be done in a functional language without the kind of variables that are used in imperative languages, without assignment statements, and

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**Figure 1.1**
The von Neumann computer architecture

![Diagram of von Neumann computer architecture](image-url)
1.4 Influences on Language Design

without iteration. Although many computer scientists have expounded on the myriad benefits of functional languages, such as Scheme, it is unlikely that they will displace the imperative languages until a non–von Neumann computer is designed that allows efficient execution of programs in functional languages. Among those who have bemoaned this fact, the most eloquent is John Backus (1978), the principal designer of the original version of Fortran.

In spite of the fact that the structure of imperative programming languages is modeled on a machine architecture, rather than on the abilities and inclinations of the users of programming languages, some believe that using imperative languages is somehow more natural than using a functional language. So, these people believe that even if functional programs were as efficient as imperative programs, the use of imperative programming languages would still dominate.

1.4.2 Programming Design Methodologies

The late 1960s and early 1970s brought an intense analysis, begun in large part by the structured-programming movement, of both the software development process and programming language design.

An important reason for this research was the shift in the major cost of computing from hardware to software, as hardware costs decreased and programmer costs increased. Increases in programmer productivity were relatively small. In addition, progressively larger and more complex problems were being solved by computers. Rather than simply solving sets of equations to simulate satellite tracks, as in the early 1960s, programs were being written for large and complex tasks, such as controlling large petroleum-refining facilities and providing worldwide airline reservation systems.

The new software development methodologies that emerged as a result of the research of the 1970s were called top-down design and stepwise refinement. The primary programming language deficiencies that were discovered were incompleteness of type checking and inadequacy of control statements (requiring the extensive use of gotos).

In the late 1970s, a shift from procedure-oriented to data-oriented program design methodologies began. Simply put, data-oriented methods emphasize data design, focusing on the use of abstract data types to solve problems.

For data abstraction to be used effectively in software system design, it must be supported by the languages used for implementation. The first language to provide even limited support for data abstraction was SIMULA 67 (Birtwistle et al., 1973), although that language certainly was not propelled to popularity because of it. The benefits of data abstraction were not widely recognized until the early 1970s. However, most languages designed since the late 1970s support data abstraction, which is discussed in detail in Chapter 11.

The latest step in the evolution of data-oriented software development, which began in the early 1980s, is object-oriented design. Object-oriented methodology begins with data abstraction, which encapsulates processing with data objects and controls access to data, and adds inheritance and dynamic
method binding. Inheritance is a powerful concept that greatly enhances the potential reuse of existing software, thereby providing the possibility of significant increases in software development productivity. This is an important factor in the increase in popularity of object-oriented languages. Dynamic (run-time) method binding allows more flexible use of inheritance.

Object-oriented programming developed along with a language that supported its concepts: Smalltalk (Goldberg and Robson, 1989). Although Smalltalk never became as widely used as many other languages, support for object-oriented programming is now part of most popular imperative languages, including Java, C++, and C#. Object-oriented concepts have also found their way into functional programming in CLOS (Bobrow et al., 1988) and F# (Syme et al., 2010), as well as logic programming in Prolog++ (Moss, 1994). Language support for object-oriented programming is discussed in detail in Chapter 12.

Procedure-oriented programming is, in a sense, the opposite of data-oriented programming. Although data-oriented methods now dominate software development, procedure-oriented methods have not been abandoned. On the contrary, in recent years, a good deal of research has occurred in procedure-oriented programming, especially in the area of concurrency. These research efforts brought with them the need for language facilities for creating and controlling concurrent program units. Java and C# include such capabilities. Concurrency is discussed in detail in Chapter 13.

All of these evolutionary steps in software development methodologies led to new language constructs to support them.

### 1.5 Language Categories

Programming languages are often categorized into four bins: imperative, functional, logic, and object oriented. However, we do not consider languages that support object-oriented programming to form a separate category of languages. We have described how the most popular languages that support object-oriented programming grew out of imperative languages. Although the object-oriented software development paradigm differs significantly from the procedure-oriented paradigm usually used with imperative languages, the extensions to an imperative language required to support object-oriented programming are not intensive. For example, the expressions, assignment statements, and control statements of C and Java are nearly identical. (On the other hand, the arrays, subprograms, and semantics of Java are very different from those of C.) Similar statements can be made for functional languages that support object-oriented programming.

Some authors refer to scripting languages as a separate category of programming languages. However, languages in this category are bound together more by their implementation method, partial or full interpretation, than by a common language design. The languages that are typically called scripting languages, among them Perl, JavaScript (Flanagan, 2011), and Ruby, are imperative languages in every sense.
A logic programming language is an example of a rule-based language. In an imperative language, an algorithm is specified in great detail, and the specific order of execution of the instructions or statements must be included. In a rule-based language, however, rules are specified in no particular order, and the language implementation system must choose an order in which the rules are used to produce the desired result. This approach to software development is radically different from those used with the other two categories of languages and clearly requires a completely different kind of language. Prolog, the most commonly used logic programming language, and logic programming are discussed in Chapter 16.

In recent years, a new category of languages has emerged, the markup/programming hybrid languages. Markup languages are not programming languages. For instance, HTML, the most widely used markup language, is used to specify the layout of information in Web documents. However, some programming capability has crept into some extensions to HTML and XML. Among these are the Java Server Pages Standard Tag Library (JSTL) and eXtensible Stylesheet Language Transformations (XSLT). Both of these are briefly introduced in Chapter 2. Those languages cannot be compared to any of the complete programming languages and therefore will not be discussed after Chapter 2.

1.6 Language Design Trade-Offs

The programming language evaluation criteria described in Section 1.3 provide a framework for language design. Unfortunately, that framework is self-contradictory. In his insightful paper on language design, Hoare (1973) stated that “there are so many important but conflicting criteria, that their reconciliation and satisfaction is a major engineering task.”

Two criteria that conflict are reliability and cost of execution. For example, the Java language definition demands that all references to array elements be checked to ensure that the index or indices are in their legal ranges. This step adds a great deal to the cost of execution of Java programs that contain large numbers of references to array elements. C does not require index range checking, so C programs execute faster than semantically equivalent Java programs, although Java programs are more reliable. The designers of Java traded execution efficiency for reliability.

As another example of conflicting criteria that leads directly to design trade-offs, consider the case of APL. APL includes a powerful set of operators for array operands. Because of the large number of operators, a significant number of new symbols had to be included in APL to represent the operators. Also, many APL operators can be used in a single, long, complex expression. One result of this high degree of expressivity is that, for applications involving many array operations, APL is very writable. Indeed, a huge amount of computation can be specified in a very small program. Another result is that APL programs have very poor readability. A compact and concise expression has a certain
mathematical beauty but it is difficult for anyone other than the programmer to understand. Well-known author Daniel McCracken (1970) once noted that it took him four hours to read and understand a four-line APL program. The designer of APL traded readability for writability.

The conflict between writability and reliability is a common one in language design. The pointers of C++ can be manipulated in a variety of ways, which supports highly flexible addressing of data. Because of the potential reliability problems with pointers, they are not included in Java.

Examples of conflicts among language design (and evaluation) criteria abound; some are subtle, others are obvious. It is therefore clear that the task of choosing constructs and features when designing a programming language requires many compromises and trade-offs.

1.7 Implementation Methods

As described in Section 1.4.1, two of the primary components of a computer are its internal memory and its processor. The internal memory is used to store programs and data. The processor is a collection of circuits that provides a realization of a set of primitive operations, or machine instructions, such as those for arithmetic and logic operations. In most computers, some of these instructions, which are sometimes called macroinstructions, are actually implemented with a set of instructions called microinstructions, which are defined at an even lower level. Because microinstructions are never seen by software, they will not be discussed further here.

The machine language of the computer is its set of instructions. In the absence of other supporting software, its own machine language is the only language that most hardware computers “understand.” Theoretically, a computer could be designed and built with a particular high-level language as its machine language, but it would be very complex and expensive. Furthermore, it would be highly inflexible, because it would be difficult (but not impossible) to use it with other high-level languages. The more practical machine design choice implements in hardware a very low-level language that provides the most commonly needed primitive operations and requires system software to create an interface to programs in higher-level languages.

A language implementation system cannot be the only software on a computer. Also required is a large collection of programs, called the operating system, which supplies higher-level primitives than those of the machine language. These primitives provide system resource management, input and output operations, a file management system, text and/or program editors, and a variety of other commonly needed functions. Because language implementation systems need many of the operating system facilities, they interface with the operating system rather than directly with the processor (in machine language).

The operating system and language implementations are layered over the machine language interface of a computer. These layers can be thought of as virtual computers, providing interfaces to the user at higher levels. For example,
an operating system and a C compiler provide a virtual C computer. With other compilers, a machine can become other kinds of virtual computers. Most computer systems provide several different virtual computers. User programs form another layer over the top of the layer of virtual computers. The layered view of a computer is shown in Figure 1.2.

The implementation systems of the first high-level programming languages, constructed in the late 1950s, were among the most complex software systems of that time. In the 1960s, intensive research efforts were made to understand and formalize the process of constructing these high-level language implementations. The greatest success of those efforts was in the area of syntax analysis, primarily because that part of the implementation process is an application of parts of automata theory and formal language theory that were then well understood.

## 1.7.1 Compilation

Programming languages can be implemented by any of three general methods. At one extreme, programs can be translated into machine language, which can be executed directly on the computer. This method is called a **compiler**
implementation and has the advantage of very fast program execution, once the translation process is complete. Most production implementations of languages, such as C, COBOL, and C++, are by compilers.

The language that a compiler translates is called the source language. The process of compilation and program execution takes place in several phases, the most important of which are shown in Figure 1.3.

The lexical analyzer gathers the characters of the source program into lexical units. The lexical units of a program are identifiers, special words, operators,
and punctuation symbols. The lexical analyzer ignores comments in the source program because the compiler has no use for them.

The syntax analyzer takes the lexical units from the lexical analyzer and uses them to construct hierarchical structures called **parse trees**. These parse trees represent the syntactic structure of the program. In many cases, no actual parse tree structure is constructed; rather, the information that would be required to build a tree is generated and used directly. Both lexical units and parse trees are further discussed in Chapter 3. Lexical analysis and syntax analysis, or parsing, are discussed in Chapter 4.

The intermediate code generator produces a program in a different language, at an intermediate level between the source program and the final output of the compiler: the machine language program. Intermediate languages sometimes look very much like assembly languages, and in fact, sometimes are actual assembly languages. In other cases, the intermediate code is at a level somewhat higher than an assembly language. The semantic analyzer is an integral part of the intermediate code generator. The semantic analyzer checks for errors, such as type errors, that are difficult, if not impossible, to detect during syntax analysis.

Optimization, which improves programs (usually in their intermediate code version) by making them smaller or faster or both. Because many kinds of optimization are difficult to do on machine language, most optimization is done on the intermediate code.

The code generator translates the optimized intermediate code version of the program into an equivalent machine language program.

The symbol table serves as a database for the compilation process. The primary contents of the symbol table are the type and attribute information of each user-defined name in the program. This information is placed in the symbol table by the lexical and syntax analyzers and is used by the semantic analyzer and the code generator.

As stated previously, although the machine language generated by a compiler can be executed directly on the hardware, it must nearly always be run along with some other code. Most user programs also require programs from the operating system. Among the most common of these are programs for input and output. The compiler builds calls to required system programs when they are needed by the user program. Before the machine language programs produced by a compiler can be executed, the required programs from the operating system must be found and linked to the user program. The linking operation connects the user program to the system programs by placing the addresses of the entry points of the system programs in the calls to them in the user program. The user and system code together are sometimes called a **load module**, or **executable image**. The process of collecting system programs and linking them to user programs is called **linking and loading**, or sometimes just **linking**. It is accomplished by a systems program called a **linker**.

---

4. Note that the words *program* and *code* are often used interchangeably.
In addition to systems programs, user programs must often be linked to previously compiled user programs that reside in libraries. So the linker not only links a given program to system programs, but also it may link it to other user programs.

The speed of the connection between a computer’s memory and its processor often determines the speed of the computer, because instructions often can be executed faster than they can be moved to the processor for execution. This connection is called the **von Neumann bottleneck**; it is the primary limiting factor in the speed of von Neumann architecture computers. The von Neumann bottleneck has been one of the primary motivations for the research and development of parallel computers.

### 1.7.2 Pure Interpretation

Pure interpretation lies at the opposite end (from compilation) among implementation methods. With this approach, programs are interpreted by another program called an interpreter, with no translation whatever. The interpreter program acts as a software simulation of a machine whose fetch-execute cycle deals with high-level language program statements rather than machine instructions. This software simulation obviously provides a virtual machine for the language.

Pure interpretation has the advantage of allowing easy implementation of many source-level debugging operations, because all run-time error messages can refer to source-level units. For example, if an array index is found to be out of range, the error message can easily indicate the source line of the error and the name of the array. On the other hand, this method has the serious disadvantage that execution is 10 to 100 times slower than in compiled systems. The primary source of this slowness is the decoding of the high-level language statements, which are far more complex than machine language instructions (although there may be fewer statements than instructions in equivalent machine code). Furthermore, regardless of how many times a statement is executed, it must be decoded every time. Therefore, statement decoding, rather than the connection between the processor and memory, is the bottleneck of a pure interpreter.

Another disadvantage of pure interpretation is that it often requires more space. In addition to the source program, the symbol table must be present during interpretation. Furthermore, the source program may be stored in a form designed for easy access and modification rather than one that provides for minimal size.

Although some simple early languages of the 1960s (APL, SNOBOL, and Lisp) were purely interpreted, by the 1980s, the approach was rarely used on high-level languages. However, in recent years, pure interpretation has made a significant comeback with some Web scripting languages, such as JavaScript and PHP, which are now widely used. The process of pure interpretation is shown in Figure 1.4.
1.7.3 Hybrid Implementation Systems

Some language implementation systems are a compromise between compilers and pure interpreters; they translate high-level language programs to an intermediate language designed to allow easy interpretation. This method is faster than pure interpretation because the source language statements are decoded only once. Such implementations are called **hybrid implementation systems**.

The process used in a hybrid implementation system is shown in Figure 1.5. Instead of translating intermediate language code to machine code, it simply interprets the intermediate code.

Perl is implemented with a hybrid system. Perl programs are partially compiled to detect errors before interpretation and to simplify the interpreter.

Initial implementations of Java were all hybrid. Its intermediate form, called **byte code**, provides portability to any machine that has a byte code interpreter and an associated run-time system. Together, these are called the Java Virtual Machine. There are now systems that translate Java byte code into machine code for faster execution.

A Just-in-Time (JIT) implementation system initially translates programs to an intermediate language. Then, during execution, it compiles intermediate language methods into machine code when they are called. The machine code version is kept for subsequent calls. JIT systems now are widely used for Java programs. Also, the .NET languages are all implemented with a JIT system.

Sometimes an implementor may provide both compiled and interpreted implementations for a language. In these cases, the interpreter is used to develop and debug programs. Then, after a (relatively) bug-free state is reached, the programs are compiled to increase their execution speed.
A preprocessor is a program that processes a program just before the program is compiled. Preprocessor instructions are embedded in programs. The preprocessor is essentially a macro expander. Preprocessor instructions are commonly used to specify that the code from another file is to be included. For example, the C preprocessor instruction

```
#include "myLib.h"
```

causes the preprocessor to copy the contents of `myLib.h` into the program at the position of the `#include`.

Other preprocessor instructions are used to define symbols to represent expressions. For example, one could use...
#define max(A, B) ((A) > (B) ? (A) : (B))

to determine the largest of two given expressions. For example, the expression

\[ x = \text{max}(2 \times y, z / 1.73); \]

would be expanded by the preprocessor to

\[ x = ((2 \times y) > (z / 1.73) ? (2 \times y) : (z / 1.73); \]

Notice that this is one of those cases where expression side effects can cause trouble. For example, if either of the expressions given to the `max` macro have side effects—such as `z++`—it could cause a problem. Because one of the two expression parameters is evaluated twice, this could result in `z` being incremented twice by the code produced by the macro expansion.

### 1.8 Programming Environments

A programming environment is the collection of tools used in the development of software. This collection may consist of only a file system, a text editor, a linker, and a compiler. Or it may include a large collection of integrated tools, each accessed through a uniform user interface. In the latter case, the development and maintenance of software is greatly enhanced. Therefore, the characteristics of a programming language are not the only measure of the software development capability of a system. We now briefly describe several programming environments.

**UNIX** is an older programming environment, first distributed in the middle 1970s, built around a portable multiprogramming operating system. It provides a wide array of powerful support tools for software production and maintenance in a variety of languages. In the past, the most important feature absent from UNIX was a uniform interface among its tools. This made it more difficult to learn and to use. However, UNIX is now often used through a GUI that runs on top of UNIX. Examples of UNIX GUIs are the Solaris Common Desktop Environment (CDE), GNOME, and KDE. These GUIs make the interface to UNIX appear similar to that of Windows and Macintosh systems.

**Borland JBuilder** is a programming environment that provides an integrated compiler, editor, debugger, and file system for Java development, where all four are accessed through a graphical interface. JBuilder is a complex and powerful system for creating Java software.

**Microsoft Visual Studio .NET** is a relatively recent step in the evolution of software development environments. It is a large and elaborate collection of software development tools, all used through a windowed interface. This system can be used to develop software in any one of the five .NET languages: C#, Visual Basic.NET, JScript (Microsoft's version of JavaScript), F# (a functional language), and C++/CLI.

**NetBeans** is a development environment that is primarily used for Java application development but also supports JavaScript, Ruby, and PHP. Both
Visual Studio and NetBeans are more than development environments—they are also frameworks, which means they actually provide common parts of the code of the application.

**SUMMARY**

The study of programming languages is valuable for some important reasons: It increases our capacity to use different constructs in writing programs, enables us to choose languages for projects more intelligently, and makes learning new languages easier.

Computers are used in a wide variety of problem-solving domains. The design and evaluation of a particular programming language is highly dependent on the domain in which it is to be used.

Among the most important criteria for evaluating languages are readability, writability, reliability, and overall cost. These will be the basis on which we examine and judge the various language features discussed in the remainder of the book.

The major influences on language design have been machine architecture and software design methodologies.

Designing a programming language is primarily an engineering feat, in which a long list of trade-offs must be made among features, constructs, and capabilities.

The major methods of implementing programming languages are compilation, pure interpretation, and hybrid implementation.

Programming environments have become important parts of software development systems, in which the language is just one of the components.

**REVIEW QUESTIONS**

1. Why is it useful for a programmer to have the ability to learn new languages, even though he or she may have a good knowledge of a number of programming languages?
2. Why is it essential to choose an appropriate programming language for a specific software solution?
3. Which programming language for scientific applications was the first to be used successfully?
4. Which is the first successful high-level programming language for business?
5. In which programming language were most of the AI applications developed prior to 1990?
6. Which is the most popular markup language for Web development?
7. Why is a list of programming language evaluation criteria for the development of software controversial?
8. How does the overall simplicity of a programming language affect its readability?
9. Why is the VAX instruction design orthogonal?
10. Why does too much orthogonality cause problems?
11. Explain how “writability” is used as a measure of how easily a language can be used to create programs.
12. Why is too much orthogonality a detriment to “writability”?
13. Give an example of expressivity in a language.
14. What is type checking?
15. Give an example of how the failure to type check, at either compile time or run time, can lead to countless program errors.
16. How is the total cost of a programming language calculated?
17. What is portability of a language?
18. What is the use of the well-definedness criterion?
19. How does the execution of a machine code program on a von Neumann architecture computer occur?
20. What two programming language deficiencies were discovered as a result of the research in software development in the 1970s?
21. What are the three fundamental features of an object-oriented programming language?
22. What language was the first to support the three fundamental features of object-oriented programming?
23. What is an example of two language design criteria that are in direct conflict with each other?
24. What are the three general methods of implementing a programming language?
25. Which produces faster program execution, a compiler or a pure interpreter?
26. What role does the symbol table play in a compiler?
27. What does a linker do?
28. Why is the von Neumann bottleneck important?
29. What are the advantages in implementing a language with a pure interpreter?

1. Do you believe our capacity for abstract thought is influenced by our language skills? Support your opinion.
2. What are some features of specific programming languages you know whose rationales are a mystery to you?
3. What arguments can you make for the idea of a single language for all programming domains?
4. What arguments can you make against the idea of a single language for all programming domains?

5. Name and explain another criterion by which languages can be judged (in addition to those discussed in this chapter).

6. What common programming language statement, in your opinion, is most detrimental to readability?

7. Java uses a right brace to mark the end of all compound statements. What are the arguments for and against this design?

8. Many languages distinguish between uppercase and lowercase letters in user-defined names. What are the pros and cons of this design decision?

9. Explain the different aspects of the cost of a programming language.

10. What are the arguments for writing efficient programs even though hardware is relatively inexpensive?

11. Describe some design trade-offs between efficiency and safety in some language you know.

12. In your opinion, what major features would a perfect programming language include?

13. Was the first high-level programming language you learned implemented with a pure interpreter, a hybrid implementation system, or a compiler? (You may have to research this.)

14. Describe the advantages and disadvantages of some programming environment you have used.

15. How do type declaration statements for simple variables affect the readability of a language, considering that some languages do not require them?

16. Write an evaluation of some programming language you know, using the criteria described in this chapter.

17. Some programming languages—for example, Pascal—have used the semicolon to separate statements, while Java uses it to terminate statements. Which of these, in your opinion, is most natural and least likely to result in syntax errors? Support your answer.

18. Many contemporary languages allow two kinds of comments: one in which delimiters are used on both ends (multiple-line comments), and one in which a delimiter marks only the beginning of the comment (one-line comments). Discuss the advantages and disadvantages of each of these with respect to our criteria.
Evolution of the Major Programming Languages

2.1 Zuse’s Plankalkül
2.2 Pseudocodes
2.3 The IBM 704 and Fortran
2.4 Functional Programming: Lisp
2.5 The First Step Toward Sophistication: ALGOL 60
2.6 Computerizing Business Records: COBOL
2.7 The Beginnings of Timesharing: Basic
2.8 Everything for Everybody: PL/I
2.9 Two Early Dynamic Languages: APL and SNOBOL
2.10 The Beginnings of Data Abstraction: SIMULA 67
2.11 Orthogonal Design: ALGOL 68
2.12 Some Early Descendants of the ALGOLs
2.13 Programming Based on Logic: Prolog
2.14 History’s Largest Design Effort: Ada
2.15 Object-Oriented Programming: Smalltalk
2.16 Combining Imperative and Object-Oriented Features: C++
2.17 An Imperative-Based Object-Oriented Language: Java
2.18 Scripting Languages
2.19 The Flagship .NET Language: C#
2.20 Markup-Programming Hybrid Languages
This chapter describes the development of a collection of programming languages. It explores the environment in which each was designed and focuses on the contributions of the language and the motivation for its development. Overall language descriptions are not included; rather, we discuss only some of the new features introduced by each language. Of particular interest are the features that most influenced subsequent languages or the field of computer science.

This chapter does not include an in-depth discussion of any language feature or concept; that is left for later chapters. Brief, informal explanations of features will suffice for our trek through the development of these languages.

This chapter discusses a wide variety of languages and language concepts that will not be familiar to many readers. These topics are discussed in detail only in later chapters. Those who find this unsettling may prefer to delay reading this chapter until the rest of the book has been studied.

The choice as to which languages to discuss here was subjective, and some readers will unhappily note the absence of one or more of their favorites. However, to keep this historical coverage to a reasonable size, it was necessary to leave out some languages that some regard highly. The choices were based on our estimate of each language's importance to language development and the computing world as a whole. We also include brief discussions of some other languages that are referenced later in the book.

The organization of this chapter is as follows: The initial versions of languages generally are discussed in chronological order. However, subsequent versions of languages appear with their initial version, rather than in later sections. For example, Fortran 2003 is discussed in the section with Fortran I (1956). Also, in some cases, languages of secondary importance that are related to a language that has its own section appear in that section.

This chapter includes listings of 14 complete example programs, each in a different language. These programs are not described in this chapter; they are meant to illustrate the appearance of programs in these languages. Readers familiar with any of the common imperative languages should be able to read and understand most of the code in these programs, except those in Lisp, COBOL, and Smalltalk. (A Scheme function similar to the Lisp example is discussed in Chapter 15.) The same problem is solved by the Fortran, ALGOL 60, PL/I, Basic, Pascal, C, Perl, Ada, Java, JavaScript, and C# programs. Note that most of the contemporary languages in this list support dynamic arrays, but because of the simplicity of the example problem, we did not use them in the example programs. Also, in the Fortran 95 program, we avoided using the features that could have avoided the use of loops altogether, in part to keep the program simple and readable and in part just to illustrate the basic loop structure of the language.

Figure 2.1 is a chart of the genealogy of the high-level languages discussed in this chapter.
Figure 2.1
Genealogy of common high-level programming languages
2.1 Zuse’s Plankalkül

The first programming language discussed in this chapter is highly unusual in several respects. For one thing, it was never implemented. Furthermore, although developed in 1945, its description was not published until 1972. Because so few people were familiar with the language, some of its capabilities did not appear in other languages until 15 years after its development.

2.1.1 Historical Background

Between 1936 and 1945, German scientist Konrad Zuse (pronounced “Tsoozuh”) built a series of complex and sophisticated computers from electromagnetic relays. By early 1945, Allied bombing had destroyed all but one of his latest models, the Z4, so he moved to a remote Bavarian village, Hinterstein, and his research group members went their separate ways.

Working alone, Zuse embarked on an effort to develop a language for expressing computations for the Z4, a project he had begun in 1943 as a proposal for his Ph.D. dissertation. He named this language Plankalkül, which means *program calculus*. In a lengthy manuscript dated 1945 but not published until 1972 (Zuse, 1972), Zuse defined Plankalkül and wrote algorithms in the language to solve a wide variety of problems.

2.1.2 Language Overview

Plankalkül was remarkably complete, with some of its most advanced features in the area of data structures. The simplest data type in Plankalkül was the single bit. Integer and floating-point numeric types were built from the bit type. The floating-point type used twos-complement notation and the “hidden bit” scheme currently used to avoid storing the most significant bit of the normalized fraction part of a floating-point value.

In addition to the usual scalar types, Plankalkül included arrays and records (called *structs* in the C-based languages). The records could include nested records.

Although the language had no explicit goto, it did include an iterative statement similar to the Ada *for*. It also had the command *Fin* with a superscript that specified an exit out of a given number of iteration loop nestings or to the beginning of a new iteration cycle. Plankalkül included a selection statement, but it did not allow an *else* clause.

One of the most interesting features of Zuse’s programs was the inclusion of mathematical expressions showing the current relationships between program variables. These expressions stated what would be true during execution at the points in the code where they appeared. These are very similar to the assertions of Java and in those in axiomatic semantics, which is discussed in Chapter 3.
Zuse’s manuscript contained programs of far greater complexity than any written prior to 1945. Included were programs to sort arrays of numbers; test the connectivity of a given graph; carry out integer and floating-point operations, including square root; and perform syntax analysis on logic formulas that had parentheses and operators in six different levels of precedence. Perhaps most remarkable were his 49 pages of algorithms for playing chess, a game in which he was not an expert.

If a computer scientist had found Zuse’s description of Plankalkül in the early 1950s, the single aspect of the language that would have hindered its implementation as defined would have been the notation. Each statement consisted of either two or three lines of code. The first line was most like the statements of current languages. The second line, which was optional, contained the subscripts of the array references in the first line. The same method of indicating subscripts was used by Charles Babbage in programs for his Analytical Engine in the middle of the nineteenth century. The last line of each Plankalkül statement contained the type names for the variables mentioned in the first line. This notation is quite intimidating when first seen.

The following example assignment statement, which assigns the value of the expression \( A[4] + 1 \) to \( A[5] \), illustrates this notation. The row labeled \( V \) is for subscripts, and the row labeled \( S \) is for the data types. In this example, \( 1.n \) means an integer of \( n \) bits:

\[
\begin{array}{c|c}
| & \text{A} + 1 => \\
V & 4 \\
S & 1.n \\
\end{array}
\]

We can only speculate on the direction that programming language design might have taken if Zuse’s work had been widely known in 1945 or even 1950. It is also interesting to consider how his work might have been different had he done it in a peaceful environment surrounded by other scientists, rather than in Germany in 1945 in virtual isolation.

## 2.2 Pseudocodes

First, note that the word *pseudocode* is used here in a different sense than its contemporary meaning. We call the languages discussed in this section pseudocodes because that’s what they were named at the time they were developed and used (the late 1940s and early 1950s). However, they are clearly not pseudocodes in the contemporary sense.

The computers that became available in the late 1940s and early 1950s were far less usable than those of today. In addition to being slow, unreliable, expensive, and having extremely small memories, the machines of that time were difficult to program because of the lack of supporting software.

There were no high-level programming languages or even assembly languages, so programming was done in machine code, which is both tedious and
error prone. Among its problems is the use of numeric codes for specifying instructions. For example, an ADD instruction might be specified by the code 14 rather than a connotative textual name, even if only a single letter. This makes programs very difficult to read. A more serious problem is absolute addressing, which makes program modification tedious and error prone. For example, suppose we have a machine language program stored in memory. Many of the instructions in such a program refer to other locations within the program, usually to reference data or to indicate the targets of branch instructions. Inserting an instruction at any position in the program other than at the end invalidates the correctness of all instructions that refer to addresses beyond the insertion point, because those addresses must be increased to make room for the new instruction. To make the addition correctly, all instructions that refer to addresses that follow the addition must be found and modified. A similar problem occurs with deletion of an instruction. In this case, however, machine languages often include a “no operation” instruction that can replace deleted instructions, thereby avoiding the problem.

These are standard problems with all machine languages and were the primary motivations for inventing assemblers and assembly languages. In addition, most programming problems of that time were numerical and required floating-point arithmetic operations and indexing of some sort to allow the convenient use of arrays. Neither of these capabilities, however, was included in the architecture of the computers of the late 1940s and early 1950s. These deficiencies naturally led to the development of somewhat higher-level languages.

2.2.1 Short Code

The first of these new languages, named Short Code, was developed by John Mauchly in 1949 for the BINAC computer, which was one of the first successful stored-program electronic computers. Short Code was later transferred to a UNIVAC I computer (the first commercial electronic computer sold in the United States) and, for several years, was one of the primary means of programming those machines. Although little is known of the original Short Code because its complete description was never published, a programming manual for the UNIVAC I version did survive (Remington-Rand, 1952). It is safe to assume that the two versions were very similar.

The words of the UNIVAC I's memory had 72 bits, grouped as 12 six-bit bytes. Short Code consisted of coded versions of mathematical expressions that were to be evaluated. The codes were byte-pair values, and many equations could be coded in a word. The following operation codes were included:

```
01 - 06 abs value  1n (n+2)nd power
02 ) 07 +  2n (n+2)nd root
03 = 08 pause  4n if <= n
04 / 09 (  58 print and tab
```
Variables were named with byte-pair codes, as were locations to be used as constants. For example, $X_0$ and $Y_0$ could be variables. The statement

$$X_0 = \text{SQRT}(\text{ABS}(Y_0))$$

would be coded in a word as 00 $X_0$ 03 20 06 $Y_0$. The initial 00 was used as padding to fill the word. Interestingly, there was no multiplication code; multiplication was indicated by simply placing the two operands next to each other, as in algebra.

Short Code was not translated to machine code; rather, it was implemented with a pure interpreter. At the time, this process was called *automatic programming*. It clearly simplified the programming process, but at the expense of execution time. Short Code interpretation was approximately 50 times slower than machine code.

### 2.2.2 Speedcoding

In other places, interpretive systems were being developed that extended machine languages to include floating-point operations. The Speedcoding system developed by John Backus for the IBM 701 is an example of such a system (Backus, 1954). The Speedcoding interpreter effectively converted the 701 to a virtual three-address floating-point calculator. The system included pseudoinstructions for the four arithmetic operations on floating-point data, as well as operations such as square root, sine, arc tangent, exponent, and logarithm. Conditional and unconditional branches and input/output conversions were also part of the virtual architecture. To get an idea of the limitations of such systems, consider that the remaining usable memory after loading the interpreter was only 700 words and that the add instruction took 4.2 milliseconds to execute. On the other hand, Speedcoding included the novel facility of automatically incrementing address registers. This facility did not appear in hardware until the UNIVAC 1107 computers of 1962. Because of such features, matrix multiplication could be done in 12 Speedcoding instructions. Backus claimed that problems that could take two weeks to program in machine code could be programmed in a few hours using Speedcoding.

### 2.2.3 The UNIVAC “Compiling” System

Between 1951 and 1953, a team led by Grace Hopper at UNIVAC developed a series of “compiling” systems named A-0, A-1, and A-2 that expanded a pseudocode into machine code subprograms in the same way as macros are expanded into assembly language. The pseudocode source for these “compilers” was still quite primitive, although even this was a great improvement over machine code because it made source programs much shorter. Wilkes (1952) independently suggested a similar process.
2.2.4 Related Work

Other means of easing the task of programming were being developed at about the same time. At Cambridge University, David J. Wheeler (1950) developed a method of using blocks of relocatable addresses to solve, at least partially, the problem of absolute addressing, and later, Maurice V. Wilkes (also at Cambridge) extended the idea to design an assembly program that could combine chosen subroutines and allocate storage (Wilkes et al., 1951, 1957). This was indeed an important and fundamental advance.

We should also mention that assembly languages, which are quite different from the pseudocodes discussed, evolved during the early 1950s. However, they had little impact on the design of high-level languages.

2.3 The IBM 704 and Fortran

Certainly one of the greatest single advances in computing came with the introduction of the IBM 704 in 1954, in large measure because its capabilities prompted the development of Fortran. One could argue that if it had not been IBM with the 704 and Fortran, it would soon thereafter have been some other organization with a similar computer and related high-level language. However, IBM was the first with both the foresight and the resources to undertake these developments.

2.3.1 Historical Background

One of the primary reasons why the slowness of interpretive systems was tolerated from the late 1940s to the mid-1950s was the lack of floating-point hardware in the available computers. All floating-point operations had to be simulated in software, a very time-consuming process. Because so much processor time was spent in software floating-point processing, the overhead of interpretation and the simulation of indexing were relatively insignificant. As long as floating-point had to be done by software, interpretation was an acceptable expense. However, many programmers of that time never used interpretive systems, preferring the efficiency of hand-coded machine (or assembly) language. The announcement of the IBM 704 system, with both indexing and floating-point instructions in hardware, heralded the end of the interpretive era, at least for scientific computation. The inclusion of floating-point hardware removed the hiding place for the cost of interpretation.

Although Fortran is often credited with being the first compiled high-level language, the question of who deserves credit for implementing the first such language is somewhat open. Knuth and Pardo (1977) give the credit to Alick E. Glennie for his Autocode compiler for the Manchester Mark I computer. Glennie developed the compiler at Fort Halstead, Royal Armaments Research Establishment, in England. The compiler was operational by September 1952. However, according to John Backus (Wexelblat, 1981, p. 26),
Glennie’s Autocode was so low level and machine oriented that it should not
be considered a compiled system. Backus gives the credit to Laning and Zierler
at the Massachusetts Institute of Technology.

The Laning and Zierler system (Laning and Zierler, 1954) was the first
algebraic translation system to be implemented. By algebraic, we mean that it
translated arithmetic expressions, used separately coded subprograms to com-
pute transcendental functions (e.g., sine and logarithm), and included arrays.
The system was implemented on the MIT Whirlwind computer, in experi-
mental prototype form, in the summer of 1952 and in a more usable form by
May 1953. The translator generated a subroutine call to code each formula, or
expression, in the program. The source language was easy to read, and the only
actual machine instructions included were for branching. Although this work
preceded the work on Fortran, it never escaped MIT.

In spite of these earlier works, the first widely accepted compiled high-
level language was Fortran. The following subsections chronicle this important
development.

2.3.2 Design Process

Even before the 704 system was announced in May 1954, plans were begun for
Fortran. By November 1954, John Backus and his group at IBM had produced
the report titled “The IBM Mathematical FORmula TRANslating System:
FORTRAN” (IBM, 1954). This document described the first version of For-
tran, which we refer to as Fortran 0, prior to its implementation. It also boldly
stated that Fortran would provide the efficiency of hand-coded programs and
the ease of programming of the interpretive pseudocode systems. In another
burst of optimism, the document stated that Fortran would eliminate coding
errors and the debugging process. Based on this premise, the first Fortran
compiler included little syntax error checking.

The environment in which Fortran was developed was as follows: (1) Com-
puters had small memories and were slow and relatively unreliable; (2) the
primary use of computers was for scientific computations; (3) there were no
existing efficient and effective ways to program computers; and (4) because of
the high cost of computers compared to the cost of programmers, speed of
the generated object code was the primary goal of the first Fortran compilers.
The characteristics of the early versions of Fortran follow directly from this
environment.

2.3.3 Fortran I Overview

Fortran 0 was modified during the implementation period, which began in
January 1955 and continued until the release of the compiler in April 1957. The
implemented language, which we call Fortran I, is described in the first Fortran
tran I included input/output formatting, variable names of up to six characters
(it had been just two in Fortran 0), user-defined subroutines, although they
could not be separately compiled, the \texttt{If} selection statement, and the \texttt{Do} loop statement.

All of Fortran I's control statements were based on 704 instructions. It is not clear whether the 704 designers dictated the control statement design of Fortran I or whether the designers of Fortran I suggested these instructions to the 704 designers.

There were no data-typing statements in the Fortran I language. Variables whose names began with \texttt{I}, \texttt{J}, \texttt{K}, \texttt{L}, \texttt{M}, and \texttt{N} were implicitly integer type, and all others were implicitly floating-point. The choice of the letters for this convention was based on the fact that at that time scientists and engineers used letters as variable subscripts, usually \texttt{i}, \texttt{j}, and \texttt{k}. In a gesture of generosity, Fortran's designers threw in the three additional letters.

The most audacious claim made by the Fortran development group during the design of the language was that the machine code produced by the compiler would be about half as efficient as what could be produced by hand.\footnote{In fact, the Fortran team believed that the code generated by their compiler could be no less than half as fast as handwritten machine code, or the language would not be adopted by users.} This, more than anything else, made skeptics of potential users and prevented a great deal of interest in Fortran before its actual release. To almost everyone's surprise, however, the Fortran development group nearly achieved its goal in efficiency. The largest part of the 18 worker-years of effort used to construct the first compiler had been spent on optimization, and the results were remarkably effective.

The early success of Fortran is shown by the results of a survey made in April 1958. At that time, roughly half of the code being written for 704s was being written in Fortran, in spite of the skepticism of most of the programming world only a year earlier.

### 2.3.4 Fortran II

The Fortran II compiler was distributed in the spring of 1958. It fixed many of the bugs in the Fortran I compilation system and added some significant features to the language, the most important being the independent compilation of subroutines. Without independent compilation, any change in a program required that the entire program be recompiled. Fortran I's lack of independent-compilation capability, coupled with the poor reliability of the 704, placed a practical restriction on the length of programs to about 300 to 400 lines (Wexelblat, 1981, p. 68). Longer programs had a poor chance of being compiled completely before a machine failure occurred. The capability of including precompiled machine language versions of subprograms shortened the compilation process considerably and made it practical to develop much larger programs.
2.3.5 Fortrans IV, 77, 90, 95, 2003, and 2008

A Fortran III was developed, but it was never widely distributed. Fortran IV, however, became one of the most widely used programming languages of its time. It evolved over the period 1960 to 1962 and was standardized as Fortran 66 (ANSI, 1966), although that name was rarely used. Fortran IV was an improvement over Fortran II in many ways. Among its most important additions were explicit type declarations for variables, a logical if construct, and the capability of passing subprograms as parameters to other subprograms.

Fortran IV was replaced by Fortran 77, which became the new standard in 1978 (ANSI, 1978a). Fortran 77 retained most of the features of Fortran IV and added character string handling, logical loop control statements, and an if with an optional else clause.

Fortran 90 (ANSI, 1992) was dramatically different from Fortran 77. The most significant additions were dynamic arrays, records, pointers, a multiple selection statement, and modules. In addition, Fortran 90 subprograms could be recursively called.

A new concept that was included in the Fortran 90 definition was that of removing some language features from earlier versions. While Fortran 90 included all of the features of Fortran 77, the language definition included a list of constructs that were recommended for removal in the next version of the language.

Fortran 90 included two simple syntactic changes that altered the appearance of both programs and the literature describing the language. First, the required fixed format of code, which required the use of specific character positions for specific parts of statements, was dropped. For example, statement labels could appear only in the first five positions and statements could not begin before the seventh position. This rigid formatting of code was designed around the use of punch cards. The second change was that the official spelling of FORTRAN became Fortran. This change was accompanied by the change in convention of using all uppercase letters for keywords and identifiers in Fortran programs. The new convention was that only the first letter of keywords and identifiers would be uppercase.

Fortran 95 (INCITS/ISO/IEC, 1997) continued the evolution of the language, but only a few changes were made. Among other things, a new iteration construct, forall, was added to ease the task of parallelizing Fortran programs.

Fortran 2003 (Metcalf et al., 2004) added support for object-oriented programming, parameterized derived types, procedure pointers, and interoperability with the C programming language.

The latest version of Fortran, Fortran 2008 (ISO/IEC 1539-1, 2010), added support for blocks to define local scopes, co-arrays, which provide a parallel execution model, and the do concurrent construct, to specify loops without interdependencies.

2.3.6 Evaluation

The original Fortran design team thought of language design only as a necessary prelude to the critical task of designing the translator. Furthermore, it never occurred to them that Fortran would be used on computers not manufactured
by IBM. However, they were forced to consider building Fortran compilers for other IBM machines only because the successor to the 704, the 709, was announced before the 704 Fortran compiler was released. The effect that Fortran has had on the use of computers, along with the fact that all subsequent programming languages owe a debt to Fortran, is indeed impressive in light of the modest goals of its designers.

One of the features of Fortran I, and all of its successors before 90, that allows highly optimizing compilers was that the types and storage for all variables are fixed before run time. No new variables or space could be allocated during execution. This was a sacrifice of flexibility to simplicity and efficiency. It eliminated the possibility of recursive subprograms and made it difficult to implement data structures that grow or change shape dynamically. Of course, the kinds of programs that were being built at the time of the development of the early versions of Fortran were primarily numerical in nature and were simple in comparison with more recent software projects. Therefore, the sacrifice was not a great one.

The overall success of Fortran is difficult to overstate: It dramatically changed the way computers are used. This is, of course, in large part due to its being the first widely used high-level language. In comparison with concepts and languages developed later, early versions of Fortran suffer in a variety of ways, as should be expected. After all, it would not be fair to compare the performance and comfort of a 1910 Model T Ford with the performance and comfort of a 2015 Ford Mustang. Nevertheless, in spite of the inadequacies of Fortran, the momentum of the huge investment in Fortran software, among other factors, has kept it in use for nearly 60 years.

Alan Perlis, one of the designers of ALGOL 60, said of Fortran in 1978, “Fortran is the 
lingua franca of the computing world. It is the language of the streets in the best sense of the word, not in the prostitutional sense of the word. And it has survived and will survive because it has turned out to be a remarkably useful part of a very vital commerce” (Wexelblat, 1981, p. 161).

The following is an example of a Fortran 95 program:

```fortran
! Fortran 95 Example program
! Input: An integer, List_Len, where List_Len is less
! than 100, followed by List_Len-Integer values
! Output: The number of input values that are greater
! than the average of all input values
Implicit none
Integer Dimension(99) :: Int_List
Integer :: List_Len, Counter, Sum, Average, Result
Result = 0
Sum = 0
Read *, List_Len
If ((List_Len > 0) .AND. (List_Len < 100)) Then
! Read input data into an array and compute its sum
   Do Counter = 1, List_Len
      Read *, Int_List(Counter)
      Sum = Sum + Int_List(Counter)
   End Do
   Average = Sum / List_Len
   Do Counter = 1, List_Len
      If (Int_List(Counter) > Average) Then
         Result = Result + 1
      End If
   End Do
End If
```


! Compute the average  
   Average = Sum / List_Len  
! Count the values that are greater than the average  
Do Counter = 1, List_Len  
   If (Int_List(Counter) > Average) Then  
      Result = Result + 1  
   End If  
End Do  
! Print the result  
   Print *, 'Number of values > Average is:', Result  
Else  
   Print *, 'Error - list length value is not legal'  
End If  
End Program Example

2.4 Functional Programming: Lisp

The first functional programming language was invented to provide language features for list processing, the need for which grew out of the first applications in the area of artificial intelligence (AI).

2.4.1 The Beginnings of Artificial Intelligence and List Processing

Interest in AI appeared in the mid-1950s in a number of places. Some of this interest grew out of linguistics, some from psychology, and some from mathematics. Linguists were concerned with natural language processing. Psychologists were interested in modeling human information storage and retrieval, as well as other fundamental processes of the brain. Mathematicians were interested in mechanizing certain intelligent processes, such as theorem proving. All of these investigations arrived at the same conclusion: Some method must be developed to allow computers to process symbolic data in linked lists. At the time, most computation was on numeric data in arrays.

The concept of list processing was developed by Allen Newell, J. C. Shaw, and Herbert Simon at the RAND Corporation. It was first published in a classic paper that describes one of the first AI programs, the Logic Theorist, and a language in which it could be implemented (Newell and Simon, 1956). The language, named IPL-I (Information Processing Language I), was never implemented. The next version, IPL-II, was implemented on a RAND Johnniac computer. Development of IPL continued until 1960, when the description of IPL-V was published (Newell and Tonge, 1960). The low level of the IPL languages prevented their widespread use. They were actually assembly languages for a hypothetical computer, implemented with an interpreter, in which

2. Logic Theorist discovered proofs for theorems in propositional calculus.
list-processing instructions were included. Another factor that kept the IPL languages from becoming popular was their implementation on the obscure Johnniac machine.

The contributions of the IPL languages were in their list design and their demonstration that list processing was feasible and useful.

IBM became interested in AI in the mid-1950s and chose theorem proving as a demonstration area. At the time, the Fortran project was still underway. The high cost of the Fortran I compiler convinced IBM that their list processing should be attached to Fortran, rather than in the form of a new language. Thus, the Fortran list processing language (FLPL) was designed and implemented as an extension to Fortran. FLPL was used to construct a theorem prover for plane geometry, which was then considered the easiest area for mechanical theorem proving.

### 2.4.2 Lisp Design Process

John McCarthy of MIT took a summer position at the IBM Information Research Department in 1958. His goal for the summer was to investigate symbolic computations and to develop a set of requirements for doing such computations. As a pilot example problem area, he chose differentiation of algebraic expressions. From this study came a list of language requirements. Among them were the control flow methods of mathematical functions: recursion and conditional expressions. The only available high-level language of the time, Fortran I, had neither of these.

Another requirement that grew from the symbolic-differentiation investigation was the need for dynamically allocated linked lists and some kind of implicit deallocation of abandoned lists. McCarthy simply would not allow his elegant algorithm for differentiation to be cluttered with explicit deallocation statements.

Because FLPL did not support recursion, conditional expressions, dynamic storage allocation, or implicit deallocation, it was clear to McCarthy that a new language was needed.

When McCarthy returned to MIT in the fall of 1958, he and Marvin Minsky formed the MIT AI Project, with funding from the Research Laboratory for Electronics. The first important effort of the project was to produce a software system for list processing. It was to be used initially to implement a program proposed by McCarthy called the Advice Taker.3 This application became the impetus for the development of the list-processing language Lisp. The first version of Lisp is sometimes called “pure Lisp” because it is a purely functional language. In the following section, we describe the development of pure Lisp.

---

3. Advice Taker represented information with sentences written in a formal language and used a logical inferencing process to decide what to do.
2.4.3 Language Overview

2.4.3.1 Data Structures

Pure Lisp has only two kinds of data structures: atoms and lists. Atoms are either symbols, which have the form of identifiers, or numeric literals. The concept of storing symbolic information in linked lists is natural and was used in IPL-II. Such structures allow insertions and deletions at any point, operations that were then thought to be a necessary part of list processing. It was eventually determined, however, that Lisp programs rarely require these operations.

Lists are specified by delimiting their elements with parentheses. Simple lists, in which elements are restricted to atoms, have the form

\[(A \ B \ C \ D)\]

Nested list structures are also specified by parentheses. For example, the list

\[(A \ (B \ C) \ D \ (E \ (F \ G)))\]

is composed of four elements. The first is the atom \(A\); the second is the sublist \((B \ C)\); the third is the atom \(D\); the fourth is the sublist \((E \ (F \ G))\), which has as its second element the sublist \((F \ G)\).

Internally, lists are stored as single-linked list structures, in which each node has two pointers and represents a list element. A node containing an atom has its first pointer pointing to some representation of the atom, such as its symbol or numeric value, or a pointer to a sublist. A node for a sublist element has its first pointer pointing to the first node of the sublist. In both cases, the second pointer of a node points to the next element of the list. A list is referenced by a pointer to its first element.

The internal representations of the two lists shown earlier are depicted in Figure 2.2. Note that the elements of a list are shown horizontally. The last element of a list has no successor, so its link is \(\text{NIL}\), which is represented in Figure 2.2 as a diagonal line in the element. Sublists are shown with the same structure.

2.4.3.2 Processes in Functional Programming

Lisp was designed as a functional programming language. All computation in a purely functional program is accomplished by applying functions to arguments. Neither the assignment statements nor the variables that abound in imperative language programs are necessary in functional language programs. Furthermore, repetitive processes can be specified with recursive function calls, making iteration (loops) unnecessary. These basic concepts of functional programming make it significantly different from programming in an imperative language.
2.4.3.3 The Syntax of Lisp

Lisp is very different from the imperative languages, both because it is a functional programming language and because the appearance of Lisp programs is so different from those in languages like Java or C++. For example, the syntax of Java is a complicated mixture of English and algebra, while Lisp’s syntax is a model of simplicity. Program code and data have exactly the same form: parenthesized lists. Consider again the list

\[(A \ B \ C \ D)\]

When interpreted as data, it is a list of four elements. When viewed as code, it is the application of the function named \(A\) to the three parameters \(B\), \(C\), and \(D\).

2.4.4 Evaluation

Lisp completely dominated AI applications for a quarter century. Much of the cause of Lisp’s reputation for being highly inefficient has been eliminated. Many contemporary implementations are compiled, and the resulting code is much faster than running the source code on an interpreter. In addition to its success in AI, Lisp pioneered functional programming, which has proven to be a lively area of research in programming languages. As stated in Chapter 1, many programming language researchers believe functional programming is a much better approach to software development than procedural programming using imperative languages.
The following is an example of a Lisp program:

```lisp
; Lisp Example function
; The following code defines a Lisp predicate function
; that takes two lists as arguments and returns True
; if the two lists are equal, and NIL (false) otherwise
(DEFUN equal_lists (lis1 lis2)
  (COND
   ((ATOM lis1) (EQ lis1 lis2))
   ((ATOM lis2) NIL)
   ((equal_lists (CAR lis1) (CAR lis2))
    (equal_lists (CDR lis1) (CDR lis2)))
   (T NIL)
  )
)
```

### 2.4.5 Two Descendants of Lisp

Two dialects of Lisp are now widely used, Scheme and Common Lisp. These are briefly discussed in the following subsections.

#### 2.4.5.1 Scheme

The Scheme language emerged from MIT in the mid-1970s. It is characterized by its small size, its exclusive use of static scoping (discussed in Chapter 5), and its treatment of functions as first-class entities. As first-class entities, Scheme functions can be assigned to variables, passed as parameters, and returned as the values of function applications. They can also be the elements of lists. Early versions of Lisp did not provide all of these capabilities, nor did they use static scoping.

As a small language with simple syntax and semantics, Scheme is well suited to educational applications, such as courses in functional programming and general introductions to programming. Scheme is described in some detail in Chapter 15.

#### 2.4.5.2 Common Lisp

During the 1970s and early 1980s, a large number of different dialects of Lisp were developed and used. This led to the familiar problem of lack of portability among programs written in the various dialects. Common Lisp (Graham, 1996) was created in an effort to rectify this situation. Common Lisp was designed by combining the features of several dialects of Lisp developed in the early 1980s, including Scheme, into a single language. Being such an amalgam, Common Lisp is a relatively large and complex language. Its basis, however, is pure Lisp, so its syntax, primitive functions, and fundamental nature come from that language.
Recognizing the flexibility provided by dynamic scoping as well as the simplicity of static scoping, Common Lisp allows both. The default scoping for variables is static, but by declaring a variable to be special, that variable becomes dynamically scoped.

Common Lisp has a large number of data types and structures, including records, arrays, complex numbers, and character strings. It also has a form of packages for modularizing collections of functions and data providing access control.

Common Lisp is further described in Chapter 15.

### 2.4.6 Related Languages

ML (MetaLanguage; Ullman, 1998) was originally designed in the 1980s by Robin Milner at the University of Edinburgh as a metalanguage for a program verification system named Logic for Computable Functions (LCF; Milner et al., 1990). ML is primarily a functional language, but it also supports imperative programming. Unlike Lisp and Scheme, the type of every variable and expression in ML can be determined at compile time. Types are associated with objects rather than names. Types of names and expressions are inferred from their context.

Unlike Lisp and Scheme, ML does not use the parenthesized functional syntax that originated with lambda expressions. Rather, the syntax of ML resembles that of the imperative languages, such as Java and C++.

Miranda was developed by David Turner (1986) at the University of Kent in Canterbury, England, in the early 1980s. Miranda is based partly on the languages ML, SASL, and KRC. Haskell (Hudak and Fasel, 1992) is based in large part on Miranda. Like Miranda, it is a purely functional language, having no variables and no assignment statement. Another distinguishing characteristic of Haskell is its use of lazy evaluation. This means that no expression is evaluated until its value is required. This leads to some surprising capabilities in the language.

Caml (Cousineau et al., 1998) and its dialect that supports object-oriented programming, OCaml (Smith, 2006), descended from ML and Haskell. Finally, F# is a relatively new typed language based directly on OCaml. F# (Syme et al., 2010) is a .NET language with direct access to the whole .NET library. Being a .NET language also means it can smoothly interoperable with any other .NET language. F# supports both functional programming and procedural programming. It also fully supports object-oriented programming.

ML, Haskell, and F# are further discussed in Chapter 15.

### 2.5 The First Step Toward Sophistication: ALGOL 60

ALGOL 60 strongly influenced subsequent programming languages and is therefore of central importance in any historical study of languages.
2.5.1 Historical Background

ALGOL 60 was the result of efforts to design a universal programming language for scientific applications. By late 1954, the Laning and Zierler algebraic system had been in operation for over a year, and the first report on Fortran had been published. Fortran became a reality in 1957, and several other high-level languages were being developed. Most notable among them were IT, which was designed by Alan Perlis at Carnegie Tech, and two languages for the UNIVAC computers, MATH-MATIC and UNICODE. The proliferation of languages made program sharing among users difficult. Furthermore, the new languages were all growing up around single architectures, some for UNIVAC computers and some for IBM 700-series machines. In response to this blossoming of machine-dependent languages, several major computer user groups in the United States, including SHARE (the IBM scientific user group) and USE (UNIVAC Scientific Exchange, the large-scale UNIVAC scientific user group), submitted a petition to the Association for Computing Machinery (ACM) on May 10, 1957, to form a committee to study and recommend action to create a machine-independent scientific programming language. Although Fortran might have been a candidate, it could not become a universal language, because at the time it was solely owned by IBM.

Previously, in 1955, GAMM (a German acronym for Society for Applied Mathematics and Mechanics) had formed a committee to design one universal, machine-independent algorithmic language. The desire for this new language was in part due to the Europeans’ fear of being dominated by IBM. By late 1957, however, the appearance of several high-level languages in the United States convinced the GAMM subcommittee that their effort had to be widened to include the Americans, and a letter of invitation was sent to ACM. In April 1958, after Fritz Bauer of GAMM presented the formal proposal to ACM, the two groups officially agreed to a joint language design project.

2.5.2 Early Design Process

GAMM and ACM each sent four members to the first design meeting. The meeting, which was held in Zurich from May 27 to June 1, 1958, began with the following goals for the new language:

- The syntax of the language should be as close as possible to standard mathematical notation, and programs written in it should be readable with little further explanation.
- It should be possible to use the language for the description of algorithms in printed publications.
- Programs in the new language must be mechanically translatable into machine language.

The first goal indicated that the new language was to be used for scientific programming, which was the primary computer application area at that time. The second was something entirely new to the computing business. The last goal is an obvious necessity for any programming language.
The Zurich meeting succeeded in producing a language that met the stated goals, but the design process required innumerable compromises, both among individuals and between the two sides of the Atlantic. In some cases, the compromises were not so much over great issues as they were over spheres of influence. The question of whether to use a comma (the European method) or a period (the American method) for a decimal point is one example.

2.5.3 ALGOL 58 Overview

The language designed at the Zurich meeting was named the International Algorithmic Language (IAL). It was suggested during the design that the language be named ALGOL, for ALGOritmic Language, but the name was rejected because it did not reflect the international scope of the committee. During the following year, however, the name was changed to ALGOL, and the language subsequently became known as ALGOL 58.

In many ways, ALGOL 58 was a descendant of Fortran, which is quite natural. It generalized many of Fortran's features and added several new constructs and concepts. Some of the generalizations had to do with the goal of not tying the language to any particular machine, and others were attempts to make the language more flexible and powerful. A rare combination of simplicity and elegance emerged from the effort.

ALGOL 58 formalized the concept of data type, although only variables that were not floating-point required explicit declaration. It added the idea of compound statements, which most subsequent languages incorporated. Some features of Fortran that were generalized were the following: Identifiers were allowed to have any length, as opposed to Fortran I's restriction to six or fewer characters; any number of array dimensions was allowed, unlike Fortran I's limitation to no more than three; the lower bound of arrays could be specified by the programmer, whereas in Fortran it was implicitly 1; nested selection statements were allowed, which was not the case in Fortran I.

ALGOL 58 acquired the assignment operator in a rather unusual way. Zuse used the form
expression => variable
for the assignment statement in Plankalkül. Although Plankalkül had not yet been published, some of the European members of the ALGOL 58 committee were familiar with the language. The committee dabbled with the Plankalkül assignment form but, because of arguments about character set limitations, the greater-than symbol was changed to a colon. Then, largely at the insistence of the Americans, the whole statement was turned around to the Fortran form
variable := expression
The Europeans preferred the opposite form, but that would be the reverse of Fortran.

---

4. The card punches of that time did not include the greater-than symbol.
2.5.4 Reception of the ALGOL 58 Report

In December 1958, publication of the ALGOL 58 report (Perlis and Samelson, 1958) was greeted with enthusiasm. In the United States, the new language was viewed more as a collection of ideas for programming language design than as a universal standard language. Actually, the ALGOL 58 report was not meant to be a finished product but rather a preliminary document for international discussion. Nevertheless, three major design and implementation efforts used the report as their basis. At the University of Michigan, the MAD language was born (Arden et al., 1961). The U.S. Naval Electronics Group produced the NELIAC language (Huskey et al., 1963). At System Development Corporation, JOVIAL was designed and implemented (Shaw, 1963). JOVIAL, an acronym for Jules' Own Version of the International Algebraic Language, represents the only language based on ALGOL 58 to achieve widespread use (Jules was Jules I. Schwartz, one of JOVIAL's designers). JOVIAL became widely used because it was the official scientific language for the U.S. Air Force for a quarter century.

The rest of the U.S. computing community was not so kind to the new language. At first, both IBM and its major scientific user group, SHARE, seemed to embrace ALGOL 58. IBM began an implementation shortly after the report was published, and SHARE formed a subcommittee, SHARE IAL, to study the language. The subcommittee subsequently recommended that ACM standardize ALGOL 58 and that IBM implement it for all of the 700-series computers. The enthusiasm was short-lived, however. By the spring of 1959, both IBM and SHARE, through their Fortran experience, had had enough of the pain and expense of getting a new language started, both in terms of developing and using the first-generation compilers and in terms of training users in the new language and persuading them to use it. By the middle of 1959, both IBM and SHARE had developed such a vested interest in Fortran that they decided to retain it as the scientific language for the IBM 700-series machines, thereby abandoning ALGOL 58.

2.5.5 ALGOL 60 Design Process

During 1959, ALGOL 58 was furiously debated in both Europe and the United States. Large numbers of suggested modifications and additions were published in the European ALGOL Bulletin and in Communications of the ACM. One of the most important events of 1959 was the presentation of the work of the Zurich committee to the International Conference on Information Processing, for there Backus introduced his new notation for describing the syntax of programming languages, which later became known as BNF (Backus-Naur form). BNF is described in detail in Chapter 3.

In January 1960, the second ALGOL meeting was held, this time in Paris. The purpose of the meeting was to debate the 80 suggestions that had been formally submitted for consideration. Peter Naur of Denmark had become heavily involved in the development of ALGOL, even though he had not been
a member of the Zurich group. It was Naur who created and published the ALGOL Bulletin. He spent a good deal of time studying Backus's paper that introduced BNF and decided that BNF should be used to describe formally the results of the 1960 meeting. After making a few relatively minor changes to BNF, he wrote a description of the new proposed language in BNF and handed it out to the members of the 1960 group at the beginning of the meeting.

### 2.5.6 ALGOL 60 Overview

Although the 1960 meeting lasted only six days, the modifications made to ALGOL 58 were dramatic. Among the most important new developments were the following:

- The concept of block structure was introduced. This allowed the programmer to localize parts of programs by introducing new data environments, or scopes.
- Two different means of passing parameters to subprograms were allowed: pass by value and pass by name.
- Procedures were allowed to be recursive. The ALGOL 58 description was unclear on this issue. Note that although recursion was new for the imperative languages, Lisp had already provided recursive functions in 1959.
- Stack-dynamic arrays were allowed. A stack-dynamic array is one for which the subscript range or ranges are specified by variables, so that the size of the array is set at the time storage is allocated to the array, which happens when the declaration is reached during execution. Stack-dynamic arrays are described in detail in Chapter 6.

Several features that might have had a dramatic impact on the success or failure of the language were proposed and rejected. Most important among these were input and output statements with formatting, which were omitted because they were thought to be machine-dependent.

The ALGOL 60 report was published in May 1960 (Naur, 1960). A number of ambiguities still remained in the language description, and a third meeting was scheduled for April 1962 in Rome to address the problems. At this meeting the group dealt only with problems; no additions to the language were allowed. The results of this meeting were published under the title “Revised Report on the Algorithmic Language ALGOL 60” (Backus et al., 1963).

### 2.5.7 Evaluation

In some ways, ALGOL 60 was a great success; in other ways, it was a dismal failure. It succeeded in becoming, almost immediately, the only acceptable formal means of communicating algorithms in computing literature, and it remained that for more than 20 years. Every imperative programming language designed since 1960 owes something to ALGOL 60. In fact, most are direct
or indirect descendants; examples include PL/I, SIMULA 67, ALGOL 68, C, Pascal, Ada, C++, Java, and C#.

The ALGOL 58/ALGOL 60 design effort included a long list of firsts. It was the first time that an international group attempted to design a programming language. It was the first language that was designed to be machine independent. It was also the first language whose syntax was formally described. This successful use of the BNF formalism initiated several important fields of computer science: formal languages, parsing theory, and BNF-based compiler design. Finally, the structure of ALGOL 60 affected machine architecture. In the most striking example of this, an extension of the language was used as the systems language of a series of large-scale computers, the Burroughs B5000, B6000, and B7000 machines, which were designed with a hardware stack to implement efficiently the block structure and recursive subprograms of the language.

On the other side of the coin, ALGOL 60 never achieved widespread use in the United States. Even in Europe, where it was more popular than in the United States, it never became the dominant language. There are a number of reasons for its lack of acceptance. For one thing, some of the features of ALGOL 60 turned out to be too flexible; they made understanding difficult and implementation inefficient. The best example of this is the pass-by-name method of passing parameters to subprograms, which is explained in Chapter 9. The difficulties of implementing ALGOL 60 are evidenced by Rutishauser’s statement in 1967 that few, if any, implementations included the full ALGOL 60 language (Rutishauser, 1967, p. 8).

The lack of input and output statements in the language was another major reason for its lack of acceptance. Implementation-dependent input/output made programs difficult to port to other computers.

Ironically, one of the most important contributions to computer science associated with ALGOL 60, BNF, was also a factor in its lack of acceptance. Although BNF is now considered a simple and elegant means of syntax description, in 1960 it seemed strange and complicated.

Finally, although there were many other problems, the entrenchment of Fortran among users and the lack of support by IBM were probably the most important factors in ALGOL 60’s failure to gain widespread use.

The ALGOL 60 effort was never really complete, in the sense that ambiguities and obscurities were always a part of the language description (Knuth, 1967).

The following is an example of an ALGOL 60 program:

```algol
comment ALGOL 60 Example Program
Input: An integer, listlen, where listlen is less than 100, followed by listlen-integer values
Output: The number of input values that are greater than the average of all the input values;
begin
    integer array intlist [1:99];
    integer listlen, counter, sum, average, result;
```
sum := 0;
result := 0;
readint (listlen);
if (listlen > 0) ^ (listlen < 100) then
begin
comment Read input into an array and compute the average;
for counter := 1 step 1 until listlen do
begin
readint (intlist[counter]);
sum := sum + intlist[counter]
end;
comment Compute the average;
average := sum / listlen;
comment Count the input values that are > average;
for counter := 1 step 1 until listlen do
if intlist[counter] > average
then result := result + 1;
comment Print result;
printstring("The number of values > average is:");
printint (result)
else
printstring ("Error-input list length is not legal");
end

2.6 Computerizing Business Records: COBOL

The story of COBOL is, in a sense, the opposite of that of ALGOL 60. Although it has been used for 65 years, COBOL has had little effect on the design of subsequent languages, except for PL/I. It may still be the most widely used language, though it is difficult to be sure one way or the other. Perhaps the most important reason why COBOL has had little influence is that few have attempted to design a new language for business applications since it appeared. That is due in part to how well COBOL's capabilities meet the needs of its application area. Another reason is that a great deal of growth in business computing over the past 30 years has occurred in small businesses. In these businesses, very little software development has taken place. Instead, most of the software used is purchased as off-the-shelf packages for various general business applications.

5. In the late 1990s, in a study associated with the Y2K problem, it was estimated that there were approximately 800 million lines of COBOL in regular use in the 22 square miles of Manhattan.
2.6.1 Historical Background

The beginning of COBOL is somewhat similar to that of ALGOL 60, in the sense that the language was designed by a committee of people meeting for relatively short periods of time. At the time, in 1959, the state of business computing was similar to the state of scientific computing several years earlier, when Fortran was being designed. One compiled language for business applications, FLOW-MATIC, had been implemented in 1957, but it belonged to one manufacturer, UNIVAC, and was designed for that company’s computers. Another language, AIMACO, was being used by the U.S. Air Force, but it was only a minor variation of FLOW-MATIC. IBM had designed a programming language for business applications, COMTRAN (COMmercial TRANslator), but it had not yet been implemented. Several other language design projects were being planned.

2.6.2 FLOW-MATIC

The origins of FLOW-MATIC are worth at least a brief discussion, because it was the primary progenitor of COBOL. In December 1953, Grace Hopper at Remington-Rand UNIVAC wrote a proposal that was indeed prophetic. It suggested that “mathematical programs should be written in mathematical notation, data processing programs should be written in English statements” (Wexelblat, 1981, p. 16). Unfortunately, in 1953, it was impossible to convince nonprogrammers that a computer could be made to understand English words. It was not until 1955 that a similar proposal had some hope of being funded by UNIVAC management, and even then it took a prototype system to do the final convincing. Part of this selling process involved compiling and running a small program, first using English keywords, then using French keywords, and then using German keywords. This demonstration was considered remarkable by UNIVAC management and was instrumental in their acceptance of Hopper’s proposal.

2.6.3 COBOL Design Process

The first formal meeting on the subject of a common language for business applications, which was sponsored by the Department of Defense, was held at the Pentagon on May 28 and 29, 1959 (exactly one year after the Zurich ALGOL meeting). The consensus of the group was that the language, then named CBL (Common Business Language), should have the following general characteristics: Most agreed that it should use English as much as possible, although a few argued for a more mathematical notation. The language must be easy to use, even at the expense of being less powerful, in order to broaden the base of those who could program computers. In addition to making the language easy to use, it was believed that the use of English would allow managers to read programs. Finally, the design should not be overly restricted by the problems of its implementation.
One of the overriding concerns at the meeting was that steps to create this universal language should be taken quickly, as a lot of work was already being done to create other business languages. In addition to the existing languages, RCA and Sylvania were working on their own business applications languages. It was clear that the longer it took to produce a universal language, the more difficult it would be for the language to become widely used. On this basis, it was decided that there should be a quick study of existing languages. For this task, the Short Range Committee was formed.

There were early decisions to separate the statements of the language into two categories—data description and executable operations—and to have statements in these two categories be in different parts of programs. One of the debates of the Short Range Committee was over the inclusion of subscripts. Many committee members argued that subscripts were too complex for the people in data processing, who were thought to be uncomfortable with mathematical notation. Similar arguments revolved around whether arithmetic expressions should be included. The final report of the Short Range Committee, which was completed in December 1959, described the language that was later named COBOL 60.

The language specification for COBOL 60, published by the Government Printing Office in April 1960 (Department of Defense, 1960), was described as “initial.” Revised versions were published in 1961 and 1962 (Department of Defense, 1961, 1962). The language was standardized by the American National Standards Institute (ANSI) group in 1968. The next three revisions were standardized by ANSI in 1974, 1985, and 2002. The language continues to evolve today.

2.6.4 Evaluation

The COBOL language originated a number of novel concepts, some of which eventually appeared in other languages. For example, the DEFINE verb of COBOL 60 was the first high-level language construct for macros. More important, hierarchical data structures (records), which first appeared in Plankalkül, were first implemented in COBOL. They have been included in most of the imperative languages designed since then. COBOL was also the first language that allowed names to be truly connotative, because it allowed both long names (up to 30 characters) and word-connector characters (hyphens).

Overall, the data division is the strong part of COBOL’s design, whereas the procedure division is relatively weak. Every variable is defined in detail in the data division, including the number of decimal digits and the location of the implied decimal point. File records are also described with this level of detail, as are lines to be output to a printer, which makes COBOL ideal for printing accounting reports. Perhaps the most important weakness of the original procedure division was in its lack of functions. Versions of COBOL prior to the 1974 standard also did not allow subprograms with parameters.

Our final comment on COBOL: It was the first programming language whose use was mandated by the Department of Defense (DoD). This mandate
came after its initial development, because COBOL was not designed specifically for the DoD. In spite of its merits, COBOL probably would not have survived without that mandate. The poor performance of the early compilers simply made the language too expensive to use. Eventually, of course, compilers became more efficient and computers became much faster and cheaper and had much larger memories. Together, these factors allowed COBOL to succeed, inside and outside DoD. Its appearance led to the electronic mechanization of accounting, an important development by any measure.

The following is an example of a COBOL program. This program reads a file named BAL-FWD-File that contains inventory information about a certain collection of items. Among other things, each item record includes the number currently on hand (BAL-ON-HAND) and the item’s reorder point (BAL-REORDER-POINT). The reorder point is the threshold number of items on hand at which more must be ordered. The program produces a list of items that must be reordered as a file named REORDER-LISTING.

IDENTIFICATION DIVISION.
PROGRAM-ID. PRODUCE-REORDER-LISTING.

ENVIRONMENT DIVISION.
CONFIGURATION SECTION.
SOURCE-COMPUTER. DEC-VAX.
OBJECT-COMPUTER. DEC-VAX.
INPUT-OUTPUT SECTION.
FILE-CONTROL.
  SELECT BAL-FWD-FILE ASSIGN TO READER.
  SELECT REORDER-LISTING ASSIGN TO LOCAL-PRINTER.

DATA DIVISION.
FILE SECTION.
FD BAL-FWD-FILE
  LABEL RECORDS ARE STANDARD
  RECORD CONTAINS 80 CHARACTERS.

  01 BAL-FWD-CARD.
    02 BAL-ITEM-NO PICTURE IS 9(5).
    02 BAL-ITEM-DESC PICTURE IS X(20).
    02 FILLER PICTURE IS X(5).
    02 BAL-UNIT-PRICE PICTURE IS 999V99.
    02 BAL-REORDER-POINT PICTURE IS 9(5).
    02 BAL-ON-HAND PICTURE IS 9(5).
    02 BAL-ON-ORDER PICTURE IS 9(5).
    02 FILLER PICTURE IS X(30).

FD REORDER-LISTING
  LABEL RECORDS ARE STANDARD
  RECORD CONTAINS 132 CHARACTERS.

  01 REORDER-LINE.
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02 RL-ITEM-NO  PICTURE IS Z(5).
02 FILLER  PICTURE IS X(5).
02 RL-ITEM-DESC  PICTURE IS X(20).
02 FILLER  PICTURE IS X(5).
02 RL-UNIT-PRICE  PICTURE IS ZZZ.99.
02 FILLER  PICTURE IS X(5).
02 RL-AVAILABLE-STOCK  PICTURE IS Z(5).
02 FILLER  PICTURE IS X(5).
02 RL-REORDER-POINT  PICTURE IS Z(5).
02 FILLER  PICTURE IS X(71).

WORKING-STORAGE SECTION.
01 SWITCHES.
   02 CARD-EOF-SWITCH  PICTURE IS X.
01 WORK-FIELDS.
   02 AVAILABLE-STOCK  PICTURE IS 9(5).

PROCEDURE DIVISION.
000-PRODUCE-REORDER-LISTING.
   OPEN INPUT BAL-FWD-FILE.
   OPEN OUTPUT REORDER-LISTING.
   MOVE "N" TO CARD-EOF-SWITCH.
   PERFORM 100-PRODUCE-REORDER-LINE
   UNTIL CARD-EOF-SWITCH IS EQUAL TO "Y".
   CLOSE BAL-FWD-FILE.
   CLOSE REORDER-LISTING.
   STOP RUN.

100-PRODUCE-REORDER-LINE.
   PERFORM 110-READ-INVENTORY-RECORD.
   IF CARD-EOF-SWITCH IS NOT EQUAL TO "Y"
      PERFORM 120-CALCULATE-AVAILABLE-STOCK
      IF AVAILABLE-STOCK IS LESS THAN BAL-REORDER-POINT
      PERFORM 130-PRINT-REORDER-LINE.
   ENDIF.

110-READ-INVENTORY-RECORD.
   READ BAL-FWD-FILE RECORD
   AT END
      MOVE "Y" TO CARD-EOF-SWITCH.

120-CALCULATE-AVAILABLE-STOCK.
   ADD BAL-ON-HAND BAL-ON-ORDER
   GIVING AVAILABLE-STOCK.

130-PRINT-REORDER-LINE.
   MOVE SPACE TO REORDER-LINE.
   MOVE BAL-ITEM-NO TO RL-ITEM-NO.
   MOVE BAL-ITEM-DESC TO RL-ITEM-DESC.
   MOVE BAL-UNIT-PRICE TO RL-UNIT-PRICE.
Basic (Mather and Waite, 1971) is another programming language that has enjoyed widespread use but has gotten little respect. Like COBOL, it has largely been ignored by computer scientists. Also, like COBOL, in its earliest versions it was inelegant and included only a meager set of control statements.

Basic was very popular on microcomputers in the late 1970s and early 1980s. This followed directly from two of the main characteristics of early versions of Basic. It was easy for beginners to learn, especially those who were not science oriented, and its smaller dialects could be implemented on computers with very small memories. When the capabilities of microcomputers grew and other languages were implemented, the use of Basic waned. A strong resurgence in the use of Basic began with the appearance of Visual Basic (Microsoft, 1991) in the early 1990s.

### 2.7.1 Design Process

Basic (Beginner’s All-purpose Symbolic Instruction Code) was originally designed at Dartmouth College (now Dartmouth University) in New Hampshire by two mathematicians, John Kemeny and Thomas Kurtz, who, in the early 1960s, developed compilers for a variety of dialects of Fortran and ALGOL 60. Their science students generally had little trouble learning or using those languages in their studies. However, Dartmouth was primarily a liberal arts institution, where science and engineering students made up only about 25 percent of the student body. It was decided in the spring of 1963 to design a new language especially for liberal arts students. This new language would use terminals as the method of computer access. The goals of the system were as follows:

1. It must be easy for nonscience students to learn and use.
2. It must be “pleasant and friendly.”
3. It must provide fast turnaround for homework.
4. It must allow free and private access.
5. It must consider user time more important than computer time.

6. Some early microcomputers included Basic interpreters that resided in 4096 bytes of ROM.
Chapter 2  Evolution of the Major Programming Languages

The last goal was indeed a revolutionary concept. It was based at least partly on the belief that computers would become significantly cheaper as time went on, which they certainly did.

The combination of the second, third, and fourth goals led to the time-shared aspect of Basic. Only with individual access through terminals by numerous simultaneous users could these goals be met in the early 1960s.

In the summer of 1963, Kemeny began work on the compiler for the first version of Basic, using remote access to a GE 225 computer. Design and coding of the operating system for Basic began in the fall of 1963. At 4:00 a.m. on May 1, 1964, the first program using the timeshared Basic was typed in and run. In June, the number of terminals on the system grew to 11, and by the fall it had ballooned to 20.

2.7.2 Language Overview

The original version of Basic was very small and, oddly, was not interactive: There was no way for an executing program to get input data from the user. Programs were typed in, compiled, and run, in a sort of batch-oriented way. The original Basic had only 14 different statement types and a single data type—floating-point. Because it was believed that few of the targeted users would appreciate the difference between integer and floating-point types, the type was referred to as “numbers.” Overall, it was a very limited language, though quite easy to learn.

2.7.3 Evaluation

The most important aspect of the original Basic was that it was the first widely used language that was used through terminals connected to a remote computer.7 Terminals had just begun to be available at that time. Before then, most programs were entered into computers through either punched cards or paper tape.

Much of the design of Basic came from Fortran, with some minor influence from the syntax of ALGOL 60. Later, it grew in a variety of ways, with little or no effort made to standardize it. The American National Standards Institute issued a Minimal Basic standard (ANSI, 1978b), but this represented only the bare minimum of language features. In fact, the original Basic was very similar to Minimal Basic.

Although it may seem surprising, Digital Equipment Corporation used a rather elaborate version of Basic named Basic-PLUS to write significant

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7. Lisp initially was used through terminals, but it was not widely used in the early 1960s.
portions of their largest operating system for the PDP-11 minicomputers, RSTS, in the 1970s.

Basic has been criticized for the poor structure of programs written in it, among other things. By the evaluation criteria discussed in Chapter 1, specifically readability and reliability, the language does indeed fare very poorly. Clearly, the early versions of the language were not meant for and should not have been used for serious programs of any significant size. Later versions are much better suited to such tasks.

The resurgence of Basic in the 1990s was driven by the appearance of Visual Basic (VB). VB became widely used in large part because it provided a simple way of building graphical user interfaces (GUIs), hence the name Visual Basic. When .NET appeared, a new version of VB came with it, VB.NET. Although it was a significant departure from earlier versions of VB, it quickly displaced the older language. Perhaps the most important difference between VB and the .NET version is that the later version fully supports object-oriented programming.

The following is an example of a Basic program:

```
REM Basic Example Program
REM Input: An integer, listlen, where listlen is less
REM than 100, followed by listlen-integer values
REM Output: The number of input values that are greater
REM than the average of all input values
DIM intlist(99)
result = 0
sum = 0
INPUT listlen
IF listlen > 0 AND listlen < 100 THEN
REM Read input into an array and compute the sum
FOR counter = 1 TO listlen
  INPUT intlist(counter)
  sum = sum + intlist(counter)
NEXT counter
REM Compute the average
average = sum / listlen
REM Count the number of input values that are > average
FOR counter = 1 TO listlen
  IF intlist(counter) > average
    THEN result = result + 1
  NEXT counter
REM Print the result
PRINT "The number of values that are > average is:";
  result
ELSE
  PRINT "Error-input list length is not legal"
END IF
END
```
User Design and Language Design

**ALAN COOPER**

Best-selling author of *About Face: The Essentials of User Interface Design*, Alan Cooper also had a large hand in designing what can be touted as the language with the most concern for user interface design, Visual Basic. For him, it all comes down to a vision for humanizing technology.

**SOME INFORMATION ON THE BASICS**

How did you get started in all of this? I’m a high school dropout with an associate degree in programming from a California community college. My first job was as a programmer for American President Lines (one of the United States’ oldest ocean transportation companies) in San Francisco. Except for a few months here and there, I’ve remained self-employed.

What is your current job? Founder and chairman of Cooper, the company that humanizes technology (www.cooper.com).

What is or was your favorite job? Interaction design consultant.

You are very well known in the fields of language design and user interface design. Any thoughts on designing languages versus designing software, versus designing anything else? It’s pretty much the same in the world of software: Know your user.

**ABOUT THAT EARLY WINDOWS RELEASE**

In the 1980s, you started using Windows and have talked about being lured by its plusses: the graphical user interface support and the dynamically linked library that let you create tools that configured themselves. What about the parts of Windows that you eventually helped shape? I was very impressed by Microsoft’s inclusion of support for practical multitasking in Windows. This included dynamic relocation and interprocess communications. MSDOS.exe was the shell program for the first few releases of Windows. It was a terrible program, and I believed that it could be improved dramatically, and I was the guy to do it. In my spare time, I immediately began to write a better shell program than the one Windows came with. I called it Tripod. Microsoft’s original shell, MSDOS.exe, was one of the main stumbling blocks to the initial success of Windows. Tripod attempted to solve the problem by being easier to use and to configure.

When was that “Aha!” moment? It wasn’t until late in 1987, when I was interviewing a corporate client, that the key design strategy for Tripod popped into my head. As the IS manager explained to me his need to create and publish a wide range of shell solutions to his disparate user base, I realized the conundrum that there is no such thing as an ideal shell. Every user would need their own personal shell, configured to their own needs and skill levels. In an instant, I perceived the solution to the shell design problem: It would be a shell construction set; a tool where each user would be able to construct exactly the shell that he or she needed for a unique mix of applications and training.

What is so compelling about the idea of a shell that can be individualized? Instead of me telling the users what the ideal shell was, they could design their own, personalized ideal shell. With a customizable shell, a programmer would create a shell that was powerful and wide ranging but also somewhat dangerous, whereas an IT manager would create a shell that could be given to a desk clerk that exposed only those few application-specific tools that the clerk used.
“MSDOS.exe was the shell program for the first few releases of Windows. It was a terrible program, and I believed that it could be improved dramatically, and I was the guy to do it. In my spare time, I immediately began to write a better shell program than the one Windows came with.”

How did you get from writing a shell program to collaborating with Microsoft? Tripod and Ruby are the same thing. After I signed a deal with Bill Gates, I changed the name of the prototype from Tripod to Ruby. I then used the Ruby prototype as prototypes should be used: as a disposable model for constructing release-quality code. Which is what I did. MS took the release version of Ruby and added QuickBasic to it, creating VB. All of those original innovations were in Tripod/Ruby.

RUBY AS THE INCUBATOR FOR VISUAL BASIC

Let’s revisit your interest in early Windows and that DLL feature. The DLL wasn’t a thing, it was a facility in the OS. It allowed a programmer to build code objects that could be linked to at run time as opposed to only at compile time. This is what allowed me to invent the dynamically extensible parts of VB, where controls can be added by third-party vendors.

The Ruby product embodied many significant advances in software design, but two of them stand out as exceptionally successful. As I mentioned, the dynamic linking capability of Windows had always intrigued me, but having the tools and knowing what to do with them were two different things. With Ruby, I finally found two practical uses for dynamic linking, and the original program contained both. First, the language was both installable and could be extended dynamically. Second, the palette of gizmos could be added to dynamically.

Was your language in Ruby the first to have a dynamic linked library and to be linked to a visual front end? As far as I know, yes.

Using a simple example, what would this enable a programmer to do with his or her program? Purchase a control, such as a grid control, from a third-party vendor, install it on his or her computer, and have the grid control appear as an integral part of the language, including the visual programming front end.

Why do they call you “the father of Visual Basic”? Ruby came with a small language, one suited only for executing the dozen or so simple commands that a shell program needs. However, this language was implemented as a chain of DLLs, any number of which could be installed at run time. The internal parser would identify a verb and then pass it along the chain of DLLs until one of them acknowledged that it knew how to process the verb. If all of the DLLs passed, there was a syntax error. From our earliest discussions, both Microsoft and I had entertained the idea of growing the language, possibly even replacing it altogether with a “real” language. C was the candidate most frequently mentioned, but eventually, Microsoft took advantage of this dynamic interface to unplug our little shell language and replace it entirely with QuickBasic. This new marriage of language to visual front end was static and permanent, and although the original dynamic interface made the coupling possible, it was lost in the process.

SOME FINAL COMMENTS ON NEW IDEAS

In the world of programming and programming tools, including languages and environments, what projects most interest you? I’m interested in creating programming tools that are designed to help users instead of programmers.

What’s the most critical rule, famous quote, or design idea to keep in mind? Bridges are not built by engineers. They are built by ironworkers.

Similarly, software programs are not built by engineers. They are built by programmers.
PL/I represents the first large-scale attempt to design a language that could be used for a broad spectrum of application areas. All previous and most subsequent languages have focused on one particular application area, such as science, artificial intelligence, or business.

### 2.8.1 Historical Background

Like Fortran, PL/I was developed as an IBM product. By the early 1960s, the users of computers in industry had settled into two separate and quite different camps: scientific and business. From the IBM point of view, scientific programmers could use either the large-scale 7090 or the small-scale 1620 IBM computers. This group used floating-point data and arrays extensively. Fortran was the primary language, although some assembly language also was used. They had their own user group, SHARE, and had little contact with anyone who worked on business applications.

For business applications, people used the large 7080 or the small 1401 IBM computers. They needed the decimal and character string data types, as well as elaborate and efficient input and output facilities. They used COBOL, although in early 1963 when the PL/I story begins, the conversion from assembly language to COBOL was far from complete. This category of users also had its own user group, GUIDE, and seldom had contact with scientific users.

In early 1963, IBM planners perceived the beginnings of a change in this situation. The two widely separated computer user groups were moving toward each other in ways that were thought certain to create problems. Scientists began to gather large files of data to be processed. This data required more sophisticated and more efficient input and output facilities. Business applications people began to use regression analysis to build management information systems, which required floating-point data and arrays. It began to appear that computing installations would soon require two separate computers and technical staffs, supporting two very different programming languages.

These perceptions naturally led to the concept of designing a single universal computer that would be capable of doing both floating-point and decimal arithmetic, and therefore both scientific and business applications. Thus was born the concept of the IBM System/360 line of computers. Along with this came the idea of a programming language that could be used for both business and scientific applications. For good measure, features to support systems programming and list processing were thrown in. Therefore, the new language was to replace Fortran, COBOL, Lisp, and the systems applications of assembly language.

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8. At the time, large computer installations required both full-time hardware and full-time system software maintenance staff.
2.8.2 Design Process

The design effort began when IBM and SHARE formed the Advanced Language Development Committee of the SHARE Fortran Project in October 1963. This new committee quickly met and formed a subcommittee called the 3 × 3 Committee, so named because it had three members from IBM and three from SHARE. The 3 × 3 Committee met for three or four days every other week to design the language.

As with the Short Range Committee for COBOL, the initial design was scheduled for completion in a remarkably short time. Apparently, regardless of the scope of a language design effort, in the early 1960s the prevailing belief was that it could be done in three months. The first version of PL/I, which was then named Fortran VI, was supposed to be completed by December, less than three months after the committee was formed. The committee pleaded successfully on two different occasions for extensions, moving the due date back to January and then to late February 1964.

The initial design concept was that the new language would be an extension of Fortran IV, maintaining compatibility, but that goal was dropped quickly along with the name Fortran VI. Until 1965, the language was known as NPL (New Programming Language). The first published report on NPL was given at the SHARE meeting in March 1964. A more complete description followed in April, and the version that would actually be implemented was published in December 1964 (IBM, 1964) by the compiler group at the IBM Hursley Laboratory in England, which was chosen to do the implementation. In 1965, the name was changed to PL/I to avoid the confusion of the name NPL with the National Physical Laboratory in England. If the compiler had been developed outside the United Kingdom, the name might have remained NPL.

2.8.3 Language Overview

Perhaps the best single-sentence description of PL/I is that it included what were then considered the best parts of ALGOL 60 (recursion and block structure), Fortran IV (separate compilation with communication through global data), and COBOL 60 (data structures, input/output, and report-generating facilities), along with an extensive collection of new constructs, all somehow cobbled together. Because PL/I is no longer a popular language, we will not attempt, even briefly, to discuss all the features of the language, or even its most controversial constructs. Instead, we will mention some of the language’s contributions to the pool of knowledge of programming languages.

PL/I was the first programming language to have the following facilities:

- Programs were allowed to create concurrently executing subprograms. Although this was a good idea, it was poorly developed in PL/I.
- It was possible to detect and handle 23 different types of exceptions, or run-time errors.
• Subprograms were allowed to be used recursively, but the capability could be disabled, allowing more efficient linkage for nonrecursive subprograms.
• Pointers were included as a data type.
• Cross-sections of arrays could be referenced. For example, the third row of a matrix could be referenced as if it were a single-dimensioned array.

2.8.4 Evaluation

Any evaluation of PL/I must begin by recognizing the ambitiousness of the design effort. In retrospect, it appears naive to think that so many constructs could have been combined successfully. However, that judgment must be tempered by acknowledging that there was little language design experience at the time. Overall, the design of PL/I was based on the premise that any construct that was useful and could be implemented should be included, with insufficient concern about how a programmer could understand and make effective use of such a collection of constructs and features. Edsger Dijkstra, in his Turing Award Lecture (Dijkstra, 1972), made one of the strongest criticisms of the complexity of PL/I: “I absolutely fail to see how we can keep our growing programs firmly within our intellectual grip when by its sheer baroqueness the programming language—our basic tool, mind you!—already escapes our intellectual control.”

In addition to the problem with the complexity due to its large size, PL/I suffered from a number of what are now considered to be poorly designed constructs. Among these were pointers, exception handling, and concurrency, although we must point out that in all cases, these constructs had not appeared in any previous language.

In terms of usage, PL/I must be considered at least a partial success. In the 1970s, it enjoyed significant use in both business and scientific applications. It was also widely used during that time as an instructional vehicle in colleges, primarily in several subset forms, such as PL/C (Cornell, 1977) and PL/CS (Conway and Constable, 1976).

The following is an example of a PL/I program:

```pli
/* PL/I PROGRAM EXAMPLE
INPUT: AN INTEGER, LISTLEN, WHERE LISTLEN IS LESS THAN 100, FOLLOWED BY LISTLEN-INTEGER VALUES
OUTPUT: THE NUMBER OF INPUT VALUES THAT ARE GREATER THAN THE AVERAGE OF ALL INPUT VALUES */
PLIEX: PROCEDURE OPTIONS (MAIN);
DECLARE INTLIST (1:99) FIXED.
DECLARE (LISTLEN, COUNTER, SUM, AVERAGE, RESULT) FIXED;
SUM = 0;
RESULT = 0;
GET LIST (LISTLEN);
IF (LISTLEN > 0) & (LISTLEN < 100) THEN
```
The structure of this section is different from that of the other sections because the languages discussed here are very different. Neither APL nor SNOBOL had much influence on later mainstream languages. Some of the interesting features of APL are discussed later in the book.

In appearance and in purpose, APL and SNOBOL are quite different. They share two fundamental characteristics, however: dynamic typing and dynamic storage allocation. Variables in both languages are essentially untyped. A variable acquires a type when it is assigned a value, at which time it assumes the type of the value assigned. Storage is allocated to a variable only when it is assigned a value, because before that there is no way to know the amount of storage that will be needed.

### 2.9.1 Origins and Characteristics of APL

APL (Brown et al., 1988) was designed around 1960 by Kenneth E. Iverson at IBM. It was not originally designed to be an implemented programming language but rather was intended to be a vehicle for describing computer architecture.

9. However, they have had some influence on some nonmainstream languages (J is based on APL, ICON is based on SNOBOL, and AWK is partially based on SNOBOL).
APL was first described in the book from which it gets its name, *A Programming Language* (Iverson, 1962). In the mid-1960s, the first implementation of APL was developed at IBM.

APL has a large number of powerful operators that are specified with a large number of symbols, which created a problem for implementors. Initially, APL was used through IBM printing terminals. These terminals had special optional print balls that provided the odd character set required by the language. One reason APL has so many operators is that it provides a large number of unit operations on arrays. For example, the transpose of any matrix is done with a single operator. The large collection of operators provides very high expressivity but also makes APL programs difficult to read. Therefore, people think of APL as a language that is best used for “throw-away” programming. Although programs can be written quickly, they should be discarded after use because they are difficult to maintain.

APL has been around for 50 years and is still used today, although not widely. Furthermore, it has not changed a great deal over its lifetime.

### 2.9.2 Origins and Characteristics of SNOBOL

SNOBOL (pronounced “snowball”; Griswold et al., 1971) was designed in the early 1960s by three people at Bell Laboratories: D. J. Farber, R. E. Griswold, and I. P. Polonsky (Farber et al., 1964). It was designed specifically for text processing. The heart of SNOBOL is a collection of powerful operations for string pattern matching. One of the early applications of SNOBOL was for writing text editors. Because the dynamic nature of SNOBOL makes it slower than alternative languages, it is no longer used for such programs. However, SNOBOL is still a live and supported language that is used for a variety of text-processing tasks in several different application areas.

### 2.10 The Beginnings of Data Abstraction: SIMULA 67

Although SIMULA 67 never achieved widespread use and had little impact on the programmers and computing of its time, some of the constructs it introduced make it historically important.

#### 2.10.1 Design Process

Two Norwegians, Kristen Nygaard and Ole-Johan Dahl, developed the language SIMULA I between 1962 and 1964 at the Norwegian Computing Center (NCC) in Oslo. They were primarily interested in using computers for simulation but also worked in operations research. SIMULA I was designed exclusively for system simulation and was first implemented in late 1964 on a UNIVAC 1107 computer.
As soon as the SIMULA I implementation was completed, Nygaard and Dahl began efforts to extend the language by adding new features and modifying some existing constructs in order to make the language useful for general-purpose applications. The result of this work was SIMULA 67, whose design was first presented publicly in March 1967 (Dahl and Nygaard, 1967). We will discuss only SIMULA 67, although some of the features of interest in SIMULA 67 are also in SIMULA I.

2.10.2 Language Overview

SIMULA 67 is an extension of ALGOL 60, taking both block structure and the control statements from that language. The primary deficiency of ALGOL 60 (and other languages at that time) for simulation applications was the design of its subprograms. Simulation requires subprograms that are allowed to restart at the position where they previously stopped. Subprograms with this kind of control are known as coroutines because the caller and called subprograms have a somewhat equal relationship with each other, rather than the rigid master/slave relationship they have in most imperative languages.

To provide support for coroutines in SIMULA 67, the class construct was developed. This was an important development because the concept of data abstraction began with it and data abstraction provides the foundation for object-oriented programming.

It is interesting to note that the important concept of data abstraction was not developed and attributed to the class construct until 1972, when Hoare (1972) recognized the connection.

2.11 Orthogonal Design: ALGOL 68

ALGOL 68 was the source of several new ideas in language design, some of which were subsequently adopted by other languages. We include it here for that reason, even though it never achieved widespread use in either Europe or the United States.

2.11.1 Design Process

The development of the ALGOL family did not end when the revised report on ALGOL 60 appeared in 1962, although it was six years until the next design iteration was published. The resulting language, ALGOL 68 (van Wijngaarden et al., 1969), was dramatically different from its predecessor.

One of the most interesting innovations of ALGOL 68 was one of its primary design criteria: orthogonality. Recall our discussion of orthogonality in Chapter 1. The use of orthogonality resulted in several innovative features of ALGOL 68, one of which is described in the following section.
2.11.2 Language Overview

One important result of orthogonality in ALGOL 68 was its inclusion of user-defined data types. Earlier languages, such as Fortran, included only a few basic data structures. PL/I included a larger number of data structures, which made it harder to learn and difficult to implement, but it still could not provide an appropriate data structure for every need.

The approach of ALGOL 68 to data structures was to provide a few primitive types and structures and allow the user to combine those primitives to define a large number of different structures. This provision for user-defined data types was carried over to some extent into all of the major imperative languages designed since then. User-defined data types are valuable because they allow the user to design data abstractions that fit particular problems very closely. All aspects of data types are discussed in Chapter 6.

As another first in the area of data types, ALGOL 68 introduced the kind of dynamic arrays that will be termed implicit heap-dynamic in Chapter 5. A dynamic array is one in which the declaration does not specify subscript bounds. Assignments to a dynamic array cause allocation of required storage. In ALGOL 68, dynamic arrays are called flex arrays.

2.11.3 Evaluation

ALGOL 68 includes a significant number of features that had not been previously used. Its use of orthogonality, which some may argue was overdone, was nevertheless revolutionary.

ALGOL 68 repeated one of the sins of ALGOL 60, however, and it was an important factor in its limited popularity. The language was described using an elegant and concise but also unknown metalanguage. Before one could read the language-describing document (van Wijngaarden et al., 1969), he or she had to learn the new metalanguage, called van Wijngaarden grammars, which were far more complex than BNF. To make matters worse, the designers invented a collection of words to explain the grammar and the language. For example, keywords were called indicants, substring extraction was called trimming, and the process of a subprogram execution was called a coercion of deproceduring, which might be meek, firm, or something else.

It is natural to contrast the design of PL/I with that of ALGOL 68, because they appeared only a few years apart. ALGOL 68 achieved writability by the principle of orthogonality: a few primitive concepts and the unrestricted use of a few combining mechanisms. PL/I achieved writability by including a large number of fixed constructs. ALGOL 68 extended the elegant simplicity of ALGOL 60, whereas PL/I simply threw together the features of several languages to attain its goals. Of course, it must be remembered that the goal of PL/I was to provide a unified tool for a broad class of problems, whereas ALGOL 68 was targeted to a single class: scientific applications.

PL/I achieved far greater acceptance than ALGOL 68, due largely to IBM’s promotional efforts and the problems of understanding and implementing
ALGOL 68. Implementation was a difficult problem for both, but PL/I had the resources of IBM to apply to building a compiler. ALGOL 68 enjoyed no such benefactor.

2.12 Some Early Descendants of the ALGOLs

All imperative languages owe some of their design to ALGOL 60 and/or ALGOL 68. This section discusses some of the early descendants of these languages.

2.12.1 Simplicity by Design: Pascal

2.12.1.1 Historical Background

Niklaus Wirth (Wirth is pronounced “Virt”) was a member of the International Federation of Information Processing (IFIP) Working Group 2.1, which was created to continue the development of ALGOL in the mid-1960s. In August 1965, Wirth and C. A. R. (“Tony”) Hoare contributed to that effort by presenting to the group a somewhat modest proposal for additions and modifications to ALGOL 60 (Wirth and Hoare, 1966). The majority of the group rejected the proposal as being too small an improvement over ALGOL 60. Instead, a much more complex revision was developed, which eventually became ALGOL 68. Wirth, along with a few other group members, did not believe that the ALGOL 68 report should have been released, based on the complexity of both the language and the meta-language used to describe it. This position later proved to have some validity because the ALGOL 68 documents, and therefore the language, were indeed found to be challenging by the computing community.

The Wirth and Hoare version of ALGOL 60 was named ALGOL-W. It was implemented at Stanford University and was used primarily as an instructional vehicle, but only at a few universities. The primary contributions of ALGOL-W were the value-result method of passing parameters and the case statement for multiple selection. The value-result method is an alternative to ALGOL 60’s pass-by-name method. Both are discussed in Chapter 9. The case statement is discussed in Chapter 8.

Wirth’s next major design effort, again based on ALGOL 60, was his most successful: Pascal. The original published definition of Pascal appeared in 1971 (Wirth, 1971). This version was modified somewhat in the implementation process and is described in Wirth (1973). The features that are often ascribed to Pascal in fact came from earlier languages. For example, user-defined data types were introduced in ALGOL 68, the case statement in ALGOL-W, and Pascal’s records are similar to those of COBOL and PL/I.

10. Pascal is named after Blaise Pascal, a seventeenth-century French philosopher and mathematician who invented the first mechanical adding machine in 1642 (among other things).
2.12.1.2 Evaluation

The largest impact of Pascal was on the teaching of programming. In 1970, most students of computer science, engineering, and science were introduced to programming with Fortran, although some universities used PL/I, languages based on PL/I, and ALGOL-W. By the mid-1970s, Pascal had become the most widely used language for this purpose. This was quite natural, because Pascal was designed specifically for teaching programming. It was not until the late 1990s that Pascal was no longer the most commonly used language for teaching programming in colleges and universities.

Because Pascal was designed as a teaching language, it lacks several features that are essential for many kinds of applications. The best example of this is the impossibility of writing a subprogram that takes as a parameter an array of variable length. Another example is the lack of any separate compilation capability. These deficiencies naturally led to many nonstandard dialects, such as Turbo Pascal.

Pascal's popularity, for both teaching programming and other applications, was based primarily on its remarkable combination of simplicity and expressivity. Although there are some insecurities in Pascal, it is still a relatively safe language, particularly when compared with Fortran or C. By the mid-1990s, the popularity of Pascal was on the decline, both in industry and in universities, primarily due to the rise of Modula-2, Ada, and C++, all of which included features not available in Pascal.

The following is an example of a Pascal program:

```pascal
{ Pascal Example Program

Input: An integer, listlen, where listlen is less than 100, followed by listlen-integer values

Output: The number of input values that are greater than the average of all input values }

program pasex (input, output);

var
  intlisttype = array [1..99] of integer;

begin
  result := 0;
  sum := 0;
  readln (listlen);
  if ((listlen > 0) and (listlen < 100)) then
    begin
      { Read input into an array and compute the sum }
      for counter := 1 to listlen do
        begin
          readln (intlist[counter]);
          sum := sum + intlist[counter]
        end;
      { Compute the average }
      average := sum / listlen;
      for counter := 1 to listlen do
        if intlist[counter] > average then
          begin
            result := result + 1
          end;
    end;
end.
```
{ Count the number of input values that are > average }

for counter := 1 to listlen do
  if (intlist[counter] > average) then
    result := result + 1;
end

{ Print the result }

writeln ('The number of values > average is:',
  result)

else
  writeln ('Error-input list length is not legal')
end.

2.12.2 A Portable Systems Language: C

Like Pascal, C contributed little to the previously known collection of language features, but it has been widely used over a long period of time. Although originally designed for systems programming, C is well suited for a wide variety of applications.

2.12.2.1 Historical Background

C’s ancestors include CPL, BCPL, B, and ALGOL 68. CPL was developed at Cambridge University in the early 1960s. BCPL is a simple systems language, also developed at Cambridge, this time by Martin Richards in 1967 (Richards, 1969).

The first work on the UNIX operating system was done in the late 1960s by Ken Thompson at Bell Laboratories. The first version was written in assembly language. The first high-level language implemented under UNIX was B, which was based on BCPL. B was designed and implemented by Thompson in 1970.

Neither BCPL nor B is a typed language, which is an oddity among high-level languages, although both are much lower-level than a language such as Java. Being untyped means that all data are considered machine words, which, although simple, leads to many complications and insecurities. For example, there is the problem of specifying floating-point rather than integer arithmetic in an expression. In one implementation of BCPL, the variable operands of a floating-point operation were preceded by periods. Variable operands not preceded by periods were considered to be integers. An alternative to this would have been to use different symbols for the floating-point operators.

This problem, along with several others, led to the development of a new typed language based on B. Originally called NB but later named C, it was designed and implemented by Dennis Ritchie at Bell Laboratories in 1972 (Kernighan and Ritchie, 1978). In some cases through BCPL, and in other cases directly, C was influenced by ALGOL 68. This is seen in its for and switch statements, in its assigning operators, and in its treatment of pointers.
Chapter 2  Evolution of the Major Programming Languages

The only “standard” for C in its first decade and a half was the book by Kernighan and Ritchie (1978).\footnote{This language is often referred to as “K & R C.”} Over that time span, the language slowly evolved, with different implementors adding different features. In 1989, ANSI produced an official description of C (ANSI, 1989), which included many of the features that implementors had already incorporated into the language. This standard was updated in 1999 (ISO, 1999). This later version includes a few significant changes to the language. Among these are a complex data type, a Boolean data type, and C++-style comments (//). We will refer to the 1989 version, which has long been called ANSI C, as C89; we will refer to the 1999 version as C99.

2.12.2.2 Evaluation

C has adequate control statements and data-structuring facilities to allow its use in many application areas. It also has a rich set of operators that provide a high degree of expressiveness.

One of the most important reasons why C is both liked and disliked is its lack of complete type checking. For example, in versions before C99, functions could be written for which parameters were not type checked. Those who like C appreciate the flexibility; those who do not like it find it too insecure. A major reason for its great increase in popularity in the 1980s was that a compiler for it was part of the widely used UNIX operating system. This inclusion in UNIX provided an essentially free and quite good compiler that was available to programmers on many different kinds of computers.

The following is an example of a C program:

```c
/* C Example Program
Input: An integer, listlen, where listlen is less than 100, followed by listlen-integer values
Output: The number of input values that are greater than the average of all input values */

int main (){
    int intlist[99], listlen, counter, sum, average, result;
    sum = 0;
    result = 0;
    scanf("%d", &listlen);
    if ((listlen > 0) && (listlen < 100)) {
        /* Read input into an array and compute the sum */
        for (counter = 0; counter < listlen; counter++) {
            scanf("%d", &intlist[counter]);
            sum += intlist[counter];
        }
        /* Compute the average */
        average = sum / listlen;
    }
    /* Compute the average */
    average = sum / listlen;
```
/* Count the input values that are > average */
    for (counter = 0; counter < listlen; counter++)
        if (intlist[counter] > average) result++;
/* Print result */
    printf("Number of values > average is:%d\n", result);
} 
else
    printf("Error-input list length is not legal\n");

2.13 Programming Based on Logic: Prolog

Simply put, logic programming is the use of a formal logic notation to communicate computational processes to a computer. Predicate calculus is the notation used in current logic programming languages.

Programming in logic programming languages is nonprocedural. Programs in such languages do not state exactly how a result is to be computed but rather describe the necessary form and/or characteristics of the result. What is needed to provide this capability in logic programming languages is a concise means of supplying the computer with both the relevant information and an inferencing process for computing desired results. Predicate calculus supplies the basic form of communication to the computer, and the proof method, named resolution, developed first by Robinson (1965), supplies the inferencing technique.

2.13.1 Design Process

During the early 1970s, Alain Colmerauer and Phillippe Roussel in the Artificial Intelligence Group at the University of Aix-Marseille, together with Robert Kowalski of the Department of Artificial Intelligence at the University of Edinburgh, developed the fundamental design of Prolog. The primary components of Prolog are a method for specifying predicate calculus propositions and an implementation of a restricted form of resolution. Both predicate calculus and resolution are described in Chapter 16. The first Prolog interpreter was developed at Marseille in 1972. The version of the language that was implemented is described in Roussel (1975). The name Prolog is from programming logic.

2.13.2 Language Overview

Prolog programs consist of collections of statements. Prolog has only a few kinds of statements, but they can be complex.
One common use of Prolog is as a kind of intelligent database. This application provides a simple framework for discussing the Prolog language.

The database of a Prolog program consists of two kinds of statements: facts and rules. The following are examples of fact statements:

```prolog
mother(joanne, jake).
father(vern, joanne).
```

These state that joanne is the mother of jake, and vern is the father of joanne.

An example of a rule statement is

```prolog
grandparent(X, Z) :- parent(X, Y), parent(Y, Z).
```

This states that it can be deduced that X is the grandparent of Z if it is true that X is the parent of Y and Y is the parent of Z, for some specific values for the variables X, Y, and Z.

The Prolog database can be interactively queried with goal statements, an example of which is

```prolog
father(bob, darcie).
```

This asks if bob is the father of darcie. When such a query, or goal, is presented to the Prolog system, it uses its resolution process to attempt to determine the truth of the statement. If it can conclude that the goal is true, it displays “true.” If it cannot prove it, it displays “false.”

### 2.13.3 Evaluation

In the 1980s, there was a relatively small group of computer scientists who believed that logic programming provided the best hope for escaping from the complexity of imperative languages, and also from the enormous problem of producing the large amount of reliable software that was needed. So far, however, there are two major reasons why logic programming has not become more widely used. First, as with some other nonimperative approaches, programs written in logic languages thus far have proven to be highly inefficient relative to equivalent imperative programs. Second, it has been determined that it is an effective approach for only a few relatively small areas of application: certain kinds of database management systems and some areas of AI.

There is a dialect of Prolog that supports object-oriented programming: Prolog++ (Moss, 1994). Logic programming and Prolog are described in greater detail in Chapter 16.
2.14 History’s Largest Design Effort: Ada

The Ada language is the result of the most extensive and expensive language design effort ever undertaken. The following paragraphs briefly describe the evolution of Ada.

2.14.1 Historical Background

The Ada language was developed for the Department of Defense (DoD), so the state of their computing environment was instrumental in determining its form. By 1974, over half of the applications of computers in DoD were embedded systems. An embedded system is one in which the computer hardware is embedded in the device it controls or for which it provides services. Software costs were rising rapidly, primarily because of the increasing complexity of systems. More than 450 different programming languages were in use for DoD projects, and none of them was standardized by DoD. Every defense contractor could define a new and different language for every contract. Because of this language proliferation, application software was rarely reused. Furthermore, no software development tools were created (because they are usually language dependent). A great many languages were in use, but none was actually suitable for embedded systems applications. For these reasons, in 1974, the Army, Navy, and Air Force each independently proposed the development of a single high-level language for embedded systems.

2.14.2 Design Process

Noting this widespread interest, in January 1975, Malcolm Currie, director of Defense Research and Engineering, formed the High-Order Language Working Group (HOLWG), initially headed by Lt. Col. William Whitaker of the Air Force. The HOLWG had representatives from all of the military services and liaisons with Great Britain, France, and what was then West Germany. Its initial charter was to do the following:

- Identify the requirements for a new DoD high-level language.
- Evaluate existing languages to determine whether there was a viable candidate.
- Recommend adoption or implementation of a minimal set of programming languages.

In April 1975, the HOLWG produced the Strawman requirements document for the new language (Department of Defense, 1975a). This was distributed to military branches, federal agencies, selected industrial and university representatives, and interested parties in Europe.

12. This result was largely due to the widespread use of assembly language for embedded systems, along with the fact that most embedded systems used specialized processors.

After a tedious process, the many submitted proposals for the language were narrowed down to four finalists, all of which were based on Pascal. In May 1979, the Cii Honeywell/Bull language design proposal was chosen from the four finalists as the design that would be used. The Cii Honeywell/Bull design team in France, the only foreign competitor among the final four, was led by Jean Ichbiah.

In the spring of 1979, Jack Cooper of the Navy Materiel Command recommended the name for the new language, Ada, which was then adopted. The name commemorates Augusta Ada Byron (1815–1851), countess of Lovelace, mathematician, and daughter of poet Lord Byron. She is generally recognized as being the world’s first programmer. She worked with Charles Babbage on his mechanical computers, the Difference and Analytical Engines, writing programs for several numerical processes.

The design and the rationale for Ada were published by ACM in its *SIGPLAN Notices* (ACM, 1979) and distributed to a readership of more than 10,000 people. A public test and evaluation conference was held in October 1979 in Boston, with representatives from over 100 organizations from the United States and Europe. By November, more than 500 language reports had been received from 15 different countries. Most of the reports suggested small modifications rather than drastic changes and outright rejections. Based on the language reports, the next version of the requirements specification, the Stoneman document (Department of Defense, 1980a), was released in February 1980.

A revised version of the language design was completed in July 1980 and was accepted as MIL-STD 1815, the standard *Ada Language Reference Manual*. The number 1815 was chosen because it was the year of the birth of Augusta Ada Byron. Another revised version of the *Ada Language Reference Manual* was released in July 1982. In 1983, the American National Standards Institute standardized Ada. This “final” official version is described in Goos and Hartmanis (1983). The Ada language design was then frozen for a minimum of five years.

### 2.14.3 Language Overview

This subsection briefly describes four of the major contributions of the Ada language.

Packages in the Ada language provide the means for encapsulating data objects, specifications for data types, and procedures. This, in turn, provides the support for the use of data abstraction in program design.

The Ada language includes extensive facilities for exception handling, which allow the programmer to gain control after any one of a wide variety of exceptions, or run-time errors, has been detected.
Program units can be generic in Ada. For example, it is possible to write a sort procedure that uses an unspecified type for the data to be sorted. Such a generic procedure must be instantiated for a specified type before it can be used, which is done with a statement that causes the compiler to generate a version of the procedure with the given type. The availability of such generic units increases the range of program units that might be reused, rather than duplicated, by programmers.

The Ada language also provides for concurrent execution of special program units, named tasks, using the rendezvous mechanism. Rendezvous is the name of a method of intertask communication and synchronization.

2.14.4 Evaluation

Perhaps the most important aspects of the design of the Ada language to consider are the following:

- Because the design was competitive, there were no limits on participation.
- The Ada language embodies most of the concepts of software engineering and language design of the late 1970s. Although one can question the actual approaches used to incorporate these features, as well as the wisdom of including such a large number of features in a language, most agree that the features are valuable.
- Although most people did not anticipate it, the development of a compiler for the Ada language was a difficult task. Only in 1985, almost four years after the language design was completed, did truly usable Ada compilers begin to appear.

The most serious criticism of Ada in its first few years was that it was too large and too complex. In particular, Hoare (1981) stated that it should not be used for any application where reliability is critical, which is precisely the type of application for which it was designed. On the other hand, others have praised it as the epitome of language design for its time. In fact, even Hoare eventually softened his view of the language.

The following is an example of an Ada program:

```ada
-- Ada Example Program
-- Input: An integer, List_Len, where List_Len is less than 100, followed by List_Len-integer values
-- Output: The number of input values that are greater than the average of all input values
with Ada.Text_IO, Ada.Integer.Text_IO;
use Ada.Text_IO, Ada.Integer.Text_IO;
procedure Ada_Ex is
  type Int_List_Type is array (1..99) of Integer;
  Int_List : Int_List_Type;
  List_Len, Sum, Average, Result : Integer;
```
begin
  Result := 0;
  Sum := 0;
  Get (List_Len);
  if (List_Len > 0) and (List_Len < 100) then
    -- Read input data into an array and compute the sum
    for Counter := 1 .. List_Len loop
      Get (Int_List(Counter));
      Sum := Sum + Int_List(Counter);
    end loop;
    -- Compute the average
    Average := Sum / List_Len;
    -- Count the number of values that are > average
    for Counter := 1 .. List_Len loop
      if Int_List(Counter) > Average then
        Result := Result + 1;
      end if;
    end loop;
    -- Print result
    Put ("The number of values > average is:");
    Put (Result);
    New_Line;
  else
    Put_Line ("Error-input list length is not legal");
  end if;
end Ada_Ex;

2.14.5 Ada 95 and Ada 2005

Two of the most important new features of Ada 95 are described briefly in the following paragraphs. In the remainder of the book, we will use the name Ada 83 for the original version and Ada 95 (its actual name) for the later version when it is important to distinguish between the two versions. In discussions of language features common to both versions, we will use the name Ada. The Ada 95 standard language is defined in ARM (1995).

The type derivation mechanism of Ada 83 is extended in Ada 95 to allow adding new components to those inherited from a base class. This provides for inheritance, a key ingredient in object-oriented programming languages. Dynamic binding of subprogram calls to subprogram definitions is accomplished through subprogram dispatching, which is based on the tag value of derived types through classwide types. This feature provides for polymorphism, another principal feature of object-oriented programming.

The rendezvous mechanism of Ada 83 provided only a cumbersome and inefficient means of sharing data among concurrent processes. It was necessary to introduce a new task to control access to the shared data. The protected objects of Ada 95 offer an attractive alternative to this. The shared data is encapsulated in a syntactic structure that controls all access to the data, either by rendezvous or by subprogram call.
It is widely believed that the popularity of Ada 95 suffered because the Department of Defense stopped requiring its use in military software systems. There were, of course, other factors that hindered its growth in popularity. Most important among these was the widespread acceptance of C++ for object-oriented programming, which occurred before Ada 95 was released. There were several additions to Ada 95 to get Ada 2005. Among these were interfaces, similar to those of Java, more control of scheduling algorithms, and synchronized interfaces.

Ada is widely used in both commercial and defense avionics, air traffic control, and rail transportation, as well as in other areas.

2.15 Object-Oriented Programming: Smalltalk

Smalltalk was the first programming language that fully supported object-oriented programming. It is therefore an important part of any discussion of the evolution of programming languages.

2.15.1 Design Process

The concepts that led to the development of Smalltalk originated in the Ph.D. dissertation work of Alan Kay in the late 1960s at the University of Utah (Kay, 1969). Kay had the remarkable foresight to predict the future availability of powerful desktop computers. Recall that the first microcomputer systems were not marketed until the mid-1970s, and they were only remotely related to the machines envisioned by Kay, which were seen to execute a million or more instructions per second and contain several megabytes of memory. Such machines, in the form of workstations, became widely available only in the early 1980s.

Kay believed that desktop computers would be used by nonprogrammers and thus would need very powerful human-interfacing capabilities. The computers of the late 1960s were largely batch oriented and were used exclusively by professional programmers and scientists. For use by nonprogrammers, Kay determined, a computer would have to be highly interactive and use sophisticated graphics in its user interface. Some of the graphics concepts came from the LOGO experience of Seymour Papert, in which graphics were used to aid children in the use of computers (Papert, 1980).

Kay originally envisioned a system he called Dynabook, which was meant to be a general information processor. It was based in part on the Flex language, which he had helped design. Flex was based primarily on SIMULA 67. Dynabook used the paradigm of the typical desk, on which there are a number of papers, some partially covered. The top sheet is often the focus of attention, with the others temporarily out of focus. The display of Dynabook would model this scene, using screen windows to represent various sheets of paper on the desktop. The user would interact with such a display both through
keystrokes and by touching the screen with his or her fingers. After the preliminary design of Dynabook earned him a Ph.D., Kay’s goal became to see such a machine constructed.

Kay found his way to the Xerox Palo Alto Research Center (Xerox PARC) and presented his ideas on Dynabook. This led to his employment there and the subsequent birth of the Learning Research Group at Xerox. The first charge of the group was to design a language to support Kay’s programming paradigm and implement it on the best personal computer then available. These efforts resulted in an “Interim” Dynabook, consisting of a Xerox Alto workstation and Smalltalk-72 software. Together, they formed a research tool for further development. A number of research projects were conducted with this system, including several experiments to teach programming to children. Along with the experiments came further developments, leading to a sequence of languages that ended with Smalltalk-80. As the language grew, so did the power of the hardware on which it resided. By 1980, both the language and the Xerox hardware nearly matched the early vision of Alan Kay.

### 2.15.2 Language Overview

The Smalltalk world is populated by nothing but objects, from integer constants to large complex software systems. All computing in Smalltalk is done by the same uniform technique: sending a message to an object to invoke one of its methods. A reply to a message is an object, which either returns the requested information or simply notifies the sender that the requested processing has been completed. The fundamental difference between a message and a subprogram call is this: A message is sent to a data object, specifically to one of the methods defined for the object. The called method is then executed, often modifying the data of the object to which the message was sent; a subprogram call is a message to the code of a subprogram. Usually the data to be processed by the subprogram is sent to it as a parameter.\(^\text{13}\)

In Smalltalk, object abstractions are classes, which are very similar to the classes of SIMULA 67. Instances of the class can be created and are then the objects of the program.

The syntax of Smalltalk is unlike that of most other programming language, in large part because of the use of messages, rather than arithmetic and logic expressions and conventional control statements. One of the Smalltalk control constructs is illustrated in the example in the next subsection.

### 2.15.3 Evaluation

Smalltalk has done a great deal to promote two separate aspects of computing: graphical user interfaces and object-oriented programming. The windowing systems that are now the dominant method of user interfaces to software systems

\(^{13}\) Of course, a method call can also pass data to be processed by the called method.
grew out of Smalltalk. Today, the most significant software design methodologies and programming languages are object oriented. Although the origin of some of the ideas of object-oriented languages came from SIMULA 67, they reached maturation in Smalltalk. It is clear that Smalltalk’s impact on the computing world is extensive and will be long-lived.

The following is an example of a Smalltalk class definition:

```
"Smalltalk Example Program"
"The following is a class definition, instantiations of which can draw equilateral polygons of any number of sides"
class name         Polygon
superclass         Object
instance variable names  ourPen
numSides
sideLength
"Class methods"
"Create an instance"
new
  ^ super new getPen

"Get a pen for drawing polygons"
getPen
  ourPen <- Pen new defaultNib: 2

"Instance methods"
"Draw a polygon"
draw
  numSides timesRepeat: [ourPen go: sideLength;
    turn: 360 // numSides]

"Set length of sides"
length: len
sideLength <- len

"Set number of sides"
sides: num
  numSides <- num
```
2.16.1 Design Process

The first step from C toward C++ was made by Bjarne Stroustrup at Bell Laboratories in 1980. The initial modifications to C included the addition of function parameter type checking and conversion and, more significantly, classes, which are related to those of SIMULA 67 and Smalltalk. Also included were derived classes, public/private access control of inherited components, constructor and destructor methods, and friend classes. During 1981, inline functions, default parameters, and overloading of the assignment operator were added. The resulting language was called C with Classes and is described in Stroustrup (1983).

It is useful to consider some goals of C with Classes. The primary goal was to provide a language in which programs could be organized as they could be organized in SIMULA 67—that is, with classes and inheritance. A second important goal was that there should be little or no performance penalty relative to C. For example, array index range checking was not even considered because a significant performance disadvantage, relative to C, would result. A third goal of C with Classes was that it could be used for every application for which C could be used, so virtually none of the features of C would be removed, not even those considered to be unsafe.

By 1984, this language was extended by the inclusion of virtual methods, which provide dynamic binding of method calls to specific method definitions, method name and operator overloading, and reference types. This version of the language was called C++. It is described in Stroustrup (1984).

In 1985, the first available implementation appeared: a system named Cfront, which translated C++ programs into C programs. This version of Cfront and the version of C++ it implemented were named Release 1.0. It is described in Stroustrup (1986).

Between 1985 and 1989, C++ continued to evolve, based largely on user reactions to the first distributed implementation. This next version was named Release 2.0. Its Cfront implementation was released in June 1989. The most important features added to C++ Release 2.0 were support for multiple inheritance (classes with more than one parent class) and abstract classes, along with some other enhancements. Abstract classes are described in Chapter 12.

Release 3.0 of C++ evolved between 1989 and 1990. It added templates, which provide parameterized types, and exception handling. The current version of C++, which was standardized in 1998, is described in ISO (1998).

In 2002, Microsoft released its .NET computing platform, which included a new version of C++, named Managed C++, or MC++. MC++ extends C++ to provide access to the functionality of the .NET Framework. The additions include properties, delegates, interfaces, and a reference type for garbage-collected objects. Properties are discussed in Chapter 11. Delegates are briefly discussed in the introduction to C# in Section 2.19. Because .NET does not support multiple inheritance, neither does MC++.
2.16.2 Language Overview

Because C++ has both functions and methods, it supports both procedural and object-oriented programming.

Operators in C++ can be overloaded, meaning the user can create operators for existing operators on user-defined types. C++ methods can also be overloaded, meaning the user can define more than one method with the same name, provided either the numbers or types of their parameters are different.

Dynamic binding in C++ is provided by virtual methods. These methods define type-dependent operations, using overloaded methods, within a collection of classes that are related through inheritance. A pointer to an object of class A can also point to objects of classes that have class A as an ancestor. When this pointer points to an overloaded virtual method, the method of the current type is chosen dynamically.

Both methods and classes can be templated, which means that they can be parameterized. For example, a method can be written as a templated method to allow it to have versions for a variety of parameter types. Classes enjoy the same flexibility.

C++ supports multiple inheritance. The exception-handling constructs of C++ are discussed in Chapter 14.

2.16.3 Evaluation

C++ rapidly became and remains a widely used language. One factor in its popularity is the availability of good and inexpensive compilers. Another factor is that it is almost completely backward compatible with C (meaning that C programs can be, with few changes, compiled as C++ programs), and in most implementations it is possible to link C++ code with C code—and thus relatively easy for the many C programmers to learn C++. Finally, at the time C++ first appeared, when object-oriented programming began to receive widespread interest, C++ was the only available language that was suitable for large commercial software projects.

On the negative side, because C++ is a very large and complex language, it clearly suffers drawbacks similar to those of PL/I. It inherited most of the insecurities of C, which make it less safe than languages such as Ada and Java.

2.16.4 A Related Language: Objective-C

Objective-C (Kochan, 2009) is another hybrid language with both imperative and object-oriented features. Objective-C was designed by Brad Cox and Tom Love in the early 1980s. Initially, it consisted of C plus the classes and message passing of Smalltalk. Among the programming languages that were created by adding support for object-oriented programming to an imperative language, Objective-C is the only one to use the Smalltalk syntax for that support.
After Steve Jobs left Apple and founded NeXT, he licensed Objective-C and it was used to write the NeXT computer system software. NeXT also released its Objective-C compiler, along with the NeXTstep development environment and a library of utilities. After the NeXT project failed, Apple bought NeXT and used Objective-C to write MAC OS X. Objective-C is the language of all iPhone software, which explains its rapid rise in popularity after the iPhone appeared.

One characteristic that Objective-C inherited from Smalltalk is the dynamic binding of messages to objects. This means that there is no static checking of messages. If a message is sent to an object and the object cannot respond to the message, it is not known until run time, when an exception is raised.

In 2006, Apple announced Objective-C 2.0, which added a form of garbage collection and new syntax for declaring properties. Unfortunately, garbage collection is not supported by the iPhone run-time system.

Objective-C is a strict superset of C, so all of the insecurities of that language are present in Objective-C.

## 2.17 An Imperative-Based Object-Oriented Language: Java

Java’s designers started with C++, removed some constructs, changed some, and added a few others. The resulting language provides much of the power and flexibility of C++, but in a smaller, simpler, and safer language. Since that initial design, Java has grown considerably.

### 2.17.1 Design Process

Java, like many programming languages, was designed for an application for which there appeared to be no satisfactory existing language. In 1990, Sun Microsystems determined there was a need for a programming language for embedded consumer electronic devices, such as toasters, microwave ovens, and interactive TV systems. Reliability was one of the primary goals for such a language. It may not seem that reliability would be an important factor in the software for a microwave oven. If an oven had malfunctioning software, it probably would not pose a grave danger to anyone and most likely would not lead to large legal settlements. However, if the software in a particular model was found to be erroneous after a million units had been manufactured and sold, their recall would entail significant cost. Therefore, reliability is an important characteristic of the software in consumer electronic products.

After considering C and C++, it was decided that neither would be satisfactory for developing software for consumer electronic devices. Although C was relatively small, it did not provide support for object-oriented programming, which they deemed a necessity. C++ supported object-oriented programming, but it was judged to be too large and complex, in part because it also supported procedure-oriented programming. It was also believed that neither C nor C++
provided the necessary level of reliability. So, a new language, later named Java, was designed. Its design was guided by the fundamental goal of providing greater simplicity and reliability than C++ was believed to provide.

Although the initial impetus for Java was consumer electronics, none of the products with which it was used in its early years were ever marketed. Starting in 1993, when the World Wide Web became widely used, and largely because of the new graphical browsers, Java was found to be a useful tool for Web programming. In particular, Java applets, which are relatively small Java programs that are interpreted in Web browsers and whose output can be included in displayed Web documents, quickly became very popular in the middle to late 1990s. In the first few years of Java popularity, the Web was its most common application.

The Java design team was headed by James Gosling, who had previously designed the UNIX emacs editor and the NeWS windowing system.

2.17.2 Language Overview

As we stated previously, Java is based on C++ but it was specifically designed to be smaller, simpler, and more reliable. Like C++, Java has both classes and primitive types. Java arrays are instances of a predefined class, whereas in C++ they are not, although many C++ users build wrapper classes for arrays to add features like index range checking, which is implicit in Java.

Java does not have pointers, but its reference types provide some of the capabilities of pointers. These references are used to point to class instances. All objects are allocated on the heap. References are always implicitly dereferenced, when necessary. So they behave more like ordinary scalar variables.

Java has a primitive Boolean type named \texttt{boolean}, used mainly for the control expressions of its control statements (such as \texttt{if} and \texttt{while}). Unlike C and C++, arithmetic expressions cannot be used for control expressions.

One significant difference between Java and many of its predecessors that support object-oriented programming, including C++, is that it is not possible to write stand-alone subprograms in Java. All Java subprograms are methods and are defined in classes. Furthermore, methods can be called through a class or object only. One consequence of this is that while C++ supports both procedural and object-oriented programming, Java supports object-oriented programming only.

Another important difference between C++ and Java is that C++ supports multiple inheritance directly in its class definitions. Java supports only single inheritance of classes, although some of the benefits of multiple inheritance can be gained by using its interface construct.

Among the C++ constructs that were not copied into Java are structs and unions.

Java includes a relatively simple form of concurrency control through its \texttt{synchronized} modifier, which can appear on methods and blocks. In either case, it causes a lock to be attached. The lock ensures mutually exclusive access or execution. In Java, it is relatively easy to create concurrent processes, which in Java are called \texttt{threads}. 
Java uses implicit storage deallocation for its objects, often called **garbage collection**. This frees the programmer from needing to delete objects explicitly when they are no longer needed. Programs written in languages that do not have garbage collection often suffer from what is sometimes called memory leakage, which means that storage is allocated but never deallocated. This can obviously lead to eventual depletion of all available storage. Object deallocation is discussed in detail in Chapter 6.

Unlike C and C++, Java includes assignment type coercions (implicit type conversions) only if they are widening (from a “smaller” type to a “larger” type). So **int** to **float** coercions are done across the assignment operator, but **float** to **int** coercions are not.

### 2.17.3 Evaluation

The designers of Java did well at trimming the excess and/or unsafe features of C++. For example, the elimination of half of the assignment coercions that are done in C++ was clearly a step toward higher reliability. Index range checking of array accesses also makes the language safer. The addition of concurrency enhances the range of applications that can be written in the language, as do the class libraries for graphical user interfaces, database access, and networking.

Java’s portability, at least in intermediate form, has often been attributed to the design of the language, but it is not. Any language can be translated to an intermediate form and “run” on any platform that has a virtual machine for that intermediate form. The price of this kind of portability is the cost of interpretation, which traditionally has been about an order of magnitude more than execution of machine code. The initial version of the Java interpreter, called the Java Virtual Machine (JVM), indeed was at least 10 times slower than equivalent compiled C programs. However, many Java programs are now translated to machine code before being executed, using Just-in-Time (JIT) compilers. This makes the efficiency of Java programs competitive with that of programs in conventionally compiled languages such as C++, at least when array index range checking is not considered.

The use of Java increased faster than that of any other programming language. Initially, this was due to its value in programming dynamic Web documents. Clearly, one of the reasons for Java’s rapid rise to prominence is simply that programmers like its design. Some developers thought C++ was too large and complex to be practical and safe. Java offered them an alternative that has much of the power of C++, but in a simpler, safer language. Another reason is that the compiler/interpreter system for Java is free and easily obtained on the Web. Java is now widely used in a variety of different applications areas.

The most recent version of Java, Java SE8, appeared in 2014. Since the first version, significant features have been added to the language. Among these are an enumeration class, generics, a new iteration construct, lambda expressions, and numerous class libraries.
The following is an example of a Java program:

```java
// Java Example Program
// Input: An integer, listlen, where listlen is less
// than 100, followed by length-integer values
// Output: The number of input data that are greater than
// the average of all input values
import java.io.*;
class IntSort {
    public static void main(String args[]) throws IOException {
        DataInputStream in = new DataInputStream(System.in);
        int listlen,
            counter,
            sum = 0,
            average,
            result = 0;
        int[] intlist = new int[99];
        listlen = Integer.parseInt(in.readLine());
        if ((listlen > 0) && (listlen < 100)) {
            /* Read input into an array and compute the sum */
            for (counter = 0; counter < listlen; counter++) {
                intlist[counter] =
                    Integer.valueOf(in.readLine()).intValue();
                sum += intlist[counter];
            }
            /* Compute the average */
            average = sum / listlen;
            /* Count the input values that are > average */
            for (counter = 0; counter < listlen; counter++)
                if (intlist[counter] > average) result++;
            /* Print result */
            System.out.println(
                "\nNumber of values > average is:" + result);
        } //** end of then clause of if ((listlen > 0) ...
        else System.out.println(
            "Error-input list length is not legal\n");
    } //** end of method main
} //** end of class IntSort
```

2.18 Scripting Languages

Scripting languages have evolved over the past 35 years. Early scripting languages were used by putting a list of commands, called a script, in a file to be interpreted. The first of these languages, named sh (for shell), began as a small collection of commands that were interpreted as calls to system subprograms that performed utility functions, such as file management and simple file filtering. To this were added variables, control flow statements, functions, and various other capabilities, and the result is a complete programming language.
One of the most powerful and widely known of these is *ksh* (Bolsky and Korn, 1995), which was developed by David Korn at Bell Laboratories.

Another scripting language is *awk*, developed by Al Aho, Brian Kernighan, and Peter Weinberger at Bell Laboratories (Aho et al., 1988). *awk* began as a report-generation language but later became a more general-purpose language.

### 2.18.1 Origins and Characteristics of Perl

The Perl language, developed by Larry Wall, was originally a combination of *sh* and *awk*. Perl has grown significantly since its beginnings and is now a powerful, although still somewhat primitive, programming language. Although it is still often called a scripting language, it is actually more similar to a typical imperative language, since it is always compiled, at least into an intermediate language, before it is executed. Furthermore, it has all the constructs to make it applicable to a wide variety of areas of computational problems.

Perl has a number of interesting features, only a few of which are mentioned in this chapter and discussed later in the book.

Variables in Perl are statically typed and implicitly declared. There are three distinctive namespaces for variables, denoted by the first character of the variables’ names. All scalar variable names begin with dollar signs ($), all array names begin with at signs (@), and all hash names (hashes are briefly described below) begin with percent signs (%). This convention makes variable names in programs more readable than those of any other programming language.

Perl includes a large number of implicit variables. Some of them are used to store Perl parameters, such as the particular form of newline character or characters that are used in the implementation. Implicit variables are commonly used as default parameters to built-in functions and default operands for some operators. The implicit variables have distinctive—although cryptic—names, such as $! and @_. The implicit variables’ names, like the user-defined variable names, use the three namespaces, so $! is a scalar.

Perl’s arrays have two characteristics that set them apart from the arrays of the common imperative languages. First, they have dynamic length, meaning that they can grow and shrink as needed during execution. Second, arrays can be sparse, meaning that there can be gaps between the elements. These gaps do not take space in memory, and the iteration statement used for arrays, `foreach`, iterates over the missing elements.

Perl includes associative arrays, which are called *hashes*. These data structures are indexed by strings and are implicitly controlled hash tables. The Perl system supplies the hash function and increases the size of the structure when necessary.

Perl is a powerful, but somewhat dangerous, language. Its scalar type stores both strings and numbers, which are normally stored in double-precision floating-point form. Depending on the context, numbers may be coerced to strings and vice versa. If a string is used in numeric context and the string cannot be converted to a number, zero is used and there is no warning or error message
provided for the user. This can lead to errors that are not detected by the compiler or run-time system. Array indexing cannot be checked, because there is no set subscript range for any array. References to nonexistent elements return `undefined`, which is interpreted as zero in numeric context. So, there is also no error detection in array element access.

Perl’s initial use was as a UNIX utility for processing text files. It was and still is widely used as a UNIX system administration tool. When the World Wide Web appeared, Perl achieved widespread use as a common gateway interface language for use with the Web, although it is now rarely used for that purpose. Perl is used as a general-purpose language for a variety of applications, such as computational biology and artificial intelligence.

The following is an example of a Perl program:

```perl
# Perl Example Program
# Input: An integer, $listlen, where $listlen is less
#        than 100, followed by $listlen-integer values.
# Output: The number of input values that are greater than
#        the average of all input values.
($sum, $result) = (0, 0);
$listlen = <STDIN>;
if (($listlen > 0) && ($listlen < 100)) {
    # Read input into an array and compute the sum
    for ($counter = 0; $counter < $listlen; $counter++) {
        $intlist[$counter] = <STDIN>;
    }  # end of for (counter ...)
    # Compute the average
    $average = $sum / $listlen;
    # Count the input values that are > average
    foreach $num (@intlist) {
        if ($num > $average) { $result++; }
    }  # end of foreach $num ...
    # Print result
    print "Number of values > average is: $result \n";
}  # end of if (($listlen ...
else {
    print "Error--input list length is not legal \n";
}
```

### 2.18.2 Origins and Characteristics of JavaScript

Use of the Web exploded in the mid-1990s after the first graphical browsers appeared. The need for computation associated with HTML documents, which by themselves are completely static, quickly became critical. Computation on the server side was made possible with the common gateway interface (CGI), which allowed HTML documents to request the execution of programs on the server, with the results of such computations returned to the browser in the form of HTML documents. Computation on the browser end became
available with the advent of Java applets. Both of these approaches have now been replaced for the most part by newer technologies, primarily scripting languages.

JavaScript was originally developed by Brendan Eich at Netscape. Its original name was Mocha. It was later renamed LiveScript. In late 1995, LiveScript became a joint venture of Netscape and Sun Microsystems and its name was changed to JavaScript. JavaScript has gone through extensive evolution, moving from version 1.0 to version 1.5 by adding many new features and capabilities. A language standard for JavaScript was developed in the late 1990s by the European Computer Manufacturers Association (ECMA) as ECMA-262. This standard has also been approved by the International Standards Organization (ISO) as ISO-16262. Microsoft’s version of JavaScript is named JScript .NET.

Although a JavaScript interpreter could be embedded in many different applications, its most common use is embedded in Web browsers. JavaScript code is embedded in HTML documents and interpreted by the browser when the documents are displayed. The primary uses of JavaScript in Web programming are to validate form input data and create dynamic HTML documents.

In spite of its name, JavaScript is related to Java only through the use of similar syntax. Java is strongly typed, but JavaScript is dynamically typed (see Chapter 5). JavaScript’s character strings and its arrays have dynamic length. Because of this, array indices are not checked for validity, although this is required in Java. Java fully supports object-oriented programming, but JavaScript supports neither inheritance nor dynamic binding of method calls to methods.

One of the most important uses of JavaScript is for dynamically creating and modifying HTML documents. JavaScript defines an object hierarchy that matches a hierarchical model of an HTML document, which is defined by the Document Object Model. Elements of an HTML document are accessed through these objects, providing the basis for dynamic control of the elements of documents.

Following is a JavaScript script for the problem previously solved in several languages in this chapter. Note that it is assumed that this script will be called from an HTML document and interpreted by a Web browser.

```javascript
// example.js
//   Input: An integer, listLen, where listLen is less than 100, followed by listLen-numeric values
//   Output: The number of input values that are greater than the average of all input values

var intList = new Array(99);
var listLen, counter, sum = 0, result = 0;

listLen = prompt ("Please type the length of the input list", ",");
```
if ((listLen > 0) && (listLen < 100)) {

    // Get the input and compute its sum
    for (counter = 0; counter < listLen; counter++) {
        intList[counter] = prompt ("Please type the next number", "");
        sum += parseInt(intList[counter]);
    }

    // Compute the average
    average = sum / listLen;

    // Count the input values that are > average
    for (counter = 0; counter < listLen; counter++)
        if (intList[counter] > average) result++;

    // Display the results
    document.write("Number of values > average is: ",
        result, "<br />");
} else
    document.write(
        "Error - input list length is not legal <br />");

2.18.3 Origins and Characteristics of PHP

PHP (Tatroe, et al., 2013) was developed by Rasmus Lerdorf, a member of the Apache Group, in 1994. His initial motivation was to provide a tool to help track visitors to his personal Web site. In 1995, he developed a package called Personal Home Page Tools, which became the first publicly distributed version of PHP. Originally, PHP was an abbreviation for Personal Home Page. Later, its user community began using the recursive name PHP: Hypertext Preprocessor, which subsequently forced the original name into obscurity. PHP is now developed, distributed, and supported as an open-source product. PHP processors are resident on most Web servers.

PHP is an HTML-embedded server-side scripting language specifically designed for Web applications. PHP code is interpreted on the Web server when an HTML document in which it is embedded has been requested by a browser. PHP code usually produces HTML code as output, which replaces the PHP code in the HTML document. Therefore, a Web browser never sees PHP code.

PHP is similar to JavaScript in its syntactic appearance, the dynamic nature of its strings and arrays, and its use of dynamic typing. PHP’s arrays are a combination of JavaScript’s arrays and Perl’s hashes.

The original version of PHP did not support object-oriented programming, but that support was added in the second release. However, PHP does not support abstract classes or interfaces, destructors, or access controls for class members.
PHP allows simple access to HTML form data, so form processing is easy with PHP. PHP provides support for many different database management systems. This makes it a useful language for building programs that need Web access to databases.

### 2.18.4 Origins and Characteristics of Python

Python (Lutz, 2013) is a relatively recent object-oriented interpreted scripting language. Its initial design was by Guido van Rossum at Stichting Mathematisch Centrum in the Netherlands in the early 1990s. Its development is now being done by the Python Software Foundation. Python is being used for the same kinds of applications as Perl: system administration, CGI programming, and other relatively small computing tasks. Python is an open-source system and is available for most common computing platforms. The Python implementation is available at [www.python.org](http://www.python.org), which also has extensive information regarding Python.

Python’s syntax is not based directly on any commonly used language. It is type checked, but dynamically typed. Instead of arrays, Python includes three kinds of data structures: lists; immutable lists, which are called *tuples*; and hashes, which are called *dictionaries*. There is a collection of list methods, such as *append*, *insert*, *remove*, and *sort*, as well as a collection of methods for dictionaries, such as *keys*, *values*, *copy*, and *has_key*. Python also supports list comprehensions, which originated with the Haskell language. List comprehensions are discussed in Section 15.8.

Python is object oriented, includes the pattern-matching capabilities of Perl, and has exception handling. Garbage collection is used to reclaim objects when they are no longer needed.

Support for CGI programming, and form processing in particular, is provided by the *cgi* module. Modules that support cookies, networking, and database access are also available.

Python includes support for concurrency with its threads, as well as support for network programming with its sockets. It also has more support for functional programming than other nonfunctional programming languages.

One of the more interesting features of Python is that it can be easily extended by any user. The modules that support the extensions can be written in any compiled language. Extensions can add functions, variables, and object types. These extensions are implemented as additions to the Python interpreter.

### 2.18.5 Origins and Characteristics of Ruby

Ruby (Thomas et al., 2005) was designed by Yukihiro Matsumoto (aka Matz) in the early 1990s and released in 1996. Since then it has continually evolved. The motivation for Ruby was dissatisfaction of its designer with Perl and Python. Although both Perl and Python support object-oriented programming,¹⁴

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¹⁴. Actually, Python's support for object-oriented programming is partial.
neither is a pure object-oriented language, at least in the sense that each has primitive (nonobject) types and each supports functions.

The primary characterizing feature of Ruby is that it is a pure object-oriented language, just as is Smalltalk. Every data value is an object and all operations are via method calls. The operators in Ruby are only syntactic mechanisms to specify method calls for the corresponding operations. Because they are methods, they can be redefined. All classes, predefined or user defined, can be subclassed.

Both classes and objects in Ruby are dynamic in the sense that methods can be dynamically added to either. This means that both classes and objects can have different sets of methods at different times during execution. So, different instantiations of the same class can behave differently. Collections of methods, data, and constants can be included in the definition of a class.

The syntax of Ruby is related to that of Eiffel and Ada. There is no need to declare variables, because dynamic typing is used. The scope of a variable is specified in its name: A variable whose name begins with a letter has local scope; one that begins with $ is an instance variable; one that begins with $ has global scope. A number of features of Perl are present in Ruby, including implicit variables with silly names, such as $.

As is the case with Python, any user can extend and/or modify Ruby. Ruby is culturally interesting because it is the first programming language designed in Japan that has achieved relatively widespread use in the United States.

2.18.6 Origins and Characteristics of Lua

Lua was designed in the early 1990s by Roberto Ierusalimschy, Waldemar Celes, and Luís Henrique de Figueiredo at the Pontifical University of Rio de Janeiro in Brazil. It is a scripting language that supports procedural and functional programming with extensibility as one of its primary goals. Among the languages that influenced its design are Scheme, Icon, and Python.

Lua is similar to JavaScript in that it does not support object-oriented programming but it was clearly influenced by it. Both have objects that play the role of both classes and objects and both have prototype inheritance rather than class inheritance. However, in Lua, the language can be extended to support object-oriented programming.

As in Scheme, Lua's functions are first-class values. Also, Lua supports closures. These capabilities allow it to be used for functional programming. Also like Scheme, Lua has only a single data structure, although in Lua's case, it is the table. Lua’s tables extend PHP’s associate arrays, which subsume the arrays of traditional imperative languages. References to table elements can take the form of references to traditional arrays, associative arrays, or records. Because functions are first-class values, they can be stored in tables, and such tables can serve as namespaces.

15. The name Lua is derived from the Portuguese word for moon.
Lua uses garbage collection for its objects, which are all heap allocated. It uses dynamic typing, as do most of the other scripting languages. Lua is a relatively small and simple language, having only 21 reserved words. The design philosophy of the language was to provide the bare essentials and relatively simple ways to extend the language to allow it to fit a variety of application areas. Much of its extensibility derives from its table data structure, which can be customized using Lua’s metatable concept.

Lua can conveniently be used as a scripting language extension to other languages. Like early implementations of Java, Lua is translated to an intermediate code and interpreted. It easily can be embedded simply in other systems, in part because of the small size of its interpreter, which is only about 150K bytes.

### 2.19 The Flagship .NET Language: C#

C#, along with the new development platform .NET,\(^\text{16}\) was announced by Microsoft in 2000. In January 2002, production versions of both were released.

#### 2.19.1 Design Process

C# is based on C++ and Java but includes some ideas from Delphi and Visual Basic. Its lead designer, Anders Hejlsberg, also designed Turbo Pascal and Delphi, which explains the Delphi parts of the heritage of C#.

The purpose of C# is to provide a language for component-based software development, specifically for such development in the .NET Framework. In this environment, components from a variety of languages can be easily combined to form systems. All of the .NET languages, which include C#, VB.NET, Managed C++, F#, and JScript .NET,\(^\text{17}\) use the common type system (CTS). The CTS provides a common class library. All types in all five .NET languages inherit from a single class root, `System.Object`. Compilers that conform to the CTS specification create objects that can be combined into software systems. All .NET languages are compiled into the same intermediate form, Intermediate Language (IL).\(^\text{18}\) Unlike Java, however, the IL is never interpreted. A Just-in-Time compiler is used to translate IL into machine code before it is executed.

#### 2.19.2 Language Overview

Many believe that one of Java’s most important advances over C++ lies in the fact that it excludes some of C++’s features. For example, C++ supports multiple inheritance, pointers, structs, `enum` types, operator overloading, and a goto

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16. The .NET development system is briefly discussed in Chapter 1.
17. Many other languages have been modified to be .NET languages.
18. Initially, IL was called MSIL (Microsoft Intermediate Language), but apparently many people thought that name was too long.
statement, but Java includes none of these. The designers of C# obviously disagreed with this wholesale removal of features, because all of these except multiple inheritance have been included in the new language.

To the credit of C#'s designers, however, in several cases, the C# version of a C++ feature has been improved. For example, the enum types of C# are safer than those of C++, because they are never implicitly converted to integers. This allows them to be more type safe. The struct type was changed significantly, resulting in a truly useful construct, whereas in C++ it serves virtually no purpose. C#'s structs are discussed in Chapter 12. C# takes a stab at improving the switch statement that is used in C, C++, and Java. C#'s switch is discussed in Chapter 8.

Although C++ includes function pointers, they share the lack of safety that is inherent in C++'s pointers to variables. C# includes a new type, delegates, which are both object-oriented and type-safe references to subprograms. Delegates are used for implementing event handlers, controlling the execution of threads, and callbacks. Callbacks are implemented in Java with interfaces; in C++, method pointers are used.

In C#, methods can take a variable number of parameters, as long as they are all the same type. This is specified by the use of a formal parameter of array type, preceded by the params reserved word.

Both C++ and Java use two distinct typing systems: one for primitives and one for objects. In addition to being confusing, this leads to a frequent need to convert values between the two systems—for example, to put a primitive value into a collection that stores objects. C# makes the conversion between values of the two typing systems partially implicit through the implicit boxing and unboxing operations, which are discussed in detail in Chapter 12.

Among the other features of C# are rectangular arrays, which are not supported in most programming languages, and a foreach statement, which is an iterator for arrays and collection objects. A similar foreach statement is found in Perl, PHP, and Java 5.0. Also, C# includes properties, which are an alternative to public data members. Properties are specified as data members with get and set methods, which are implicitly called when references and assignments are made to the associated data members.

C# has evolved continuously and quickly from its initial release in 2002. The most recent version is C# 5.0. New in C# 5.0 are asynchronous functions.

### 2.19.3 Evaluation

C# was meant to be an improvement over both C++ and Java as a general-purpose programming language. Although it can be argued that some of its features are a step backward, C# includes some constructs that move it beyond its predecessors. Some of its features will surely be adopted by other programming languages of the future.

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19. When an object calls a method of another object and needs to be notified when that method has completed its task, the called method calls its caller back. This is known as a callback.

20. This feature was added to Java in Java 5.0.
The following is an example of a C# program:

```csharp
// C# Example Program
// Input:  An integer, listlen, where listlen is less than 100, followed by listlen-integer values.
// Output: The number of input values that are greater than the average of all input values.
using System;
public class Ch2example {
    static void Main() {
        int[] intlist;
        int listlen,
            counter,
            sum = 0,
            average,
            result = 0;
        intList = new int[99];
        listlen = Int32.Parse(Console.readLine());
        if ((listlen > 0) && (listlen < 100)) {
            // Read input into an array and compute the sum
            for (counter = 0; counter < listlen; counter++) {
                intList[counter] =
                    Int32.Parse(Console.readLine());
                sum += intList[counter];
            } // End of for (counter ...
            // Compute the average
            average = sum / listlen;
            // Count the input values that are > average
            foreach (int num in intList)
                if (num > average) result++;
            // Print result
            Console.WriteLine(
                "Number of values > average is:" + result);
        } // End of if ((listlen ...
        else
            Console.WriteLine(
                "Error--input list length is not legal");
    } // End of method Main
} // End of class Ch2example
```

### 2.20 Markup-Programming Hybrid Languages

A markup-programming hybrid language is a markup language in which some of the elements can specify programming actions, such as control flow and computation. The following subsections introduce two such hybrid languages, XSLT and JSP.
2.20.1 XSLT

eXtensible markup language (XML) is a metamarkup language. Such a lan-
guage is used to define markup languages. XML-derived markup languages are
used to define XML data documents. Although XML documents are human
readable, they are processed by computers. This processing sometimes consists
only of transformations to alternative forms that can be effectively displayed or
printed. In many cases, such transformations are to HTML, which can be dis-
played by a Web browser. In other cases, the data in the document is processed,
just as with other forms of data files.

The transformation of XML documents to HTML documents is specified
in another markup language, eXtensible stylesheet language transformations
(XSLT) (www.w3.org/TR/XSLT). XSLT can specify programming-like opera-
tions. Therefore, XSLT is a markup-programming hybrid language. XSLT was
defined by the World Wide Web Consortium (W3C) in the late 1990s.

An XSLT processor is a program that takes as input an XML data docu-
ment and an XSLT document (which is also in the form of an XML document).
In this processing, the XML data document is transformed to another XML
document, using the transformations described in the XSLT document. The
XSLT document specifies transformations by defining templates, which are
data patterns that could be found by the XSLT processor in the XML input file.
Associated with each template in the XSLT document are its transformation
instructions, which specify how the matching data is to be transformed before
being put in the output document. So, the templates (and their associated pro-
cessing) act as subprograms, which are “executed” when the XSLT processor
finds a pattern match in the data of the XML document.

XSLT also has programming constructs at a lower level. For example, a
looping construct is included, which allows repeated parts of the XML docu-
ment to be selected. There is also a sort process. These lower-level constructs
are specified with XSLT tags, such as <for-each>.

2.20.2 JSP

The “core” part of the Java Server Pages Standard Tag Library (JSTL) is
another markup-programming hybrid language, although its form and pur-
pose are different from those of XSLT. Before discussing JSTL, it is necessary
to introduce the ideas of servlets and Java Server Pages (JSP). A servlet is an
instance of a Java class that resides on and is executed on a Web server system.
The execution of a servlet is requested by a markup document being displayed
by a Web browser. The servlet’s output, which is in the form of an HTML
document, is returned to the requesting browser. A program that runs in the
Web server process, called a servlet container, controls the execution of serv-
lets. Servlets are commonly used for form processing and for database access.

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21. The output document of the XSLT processor could also be in HTML or plain text.
JSP is a collection of technologies designed to support dynamic Web documents and provide other processing needs of Web documents. When a JSP document, which is often a mixture of HTML and Java, is requested by a browser, the JSP processor program, which resides on a Web server system, converts the document to a servlet. The document’s embedded Java code is copied to the servlet. The plain HTML is copied into Java print statements that output it as is. The JSTL markup in the JSP document is processed, as discussed in the following paragraph. The servlet produced by the JSP processor is run by the servlet container.

The JSTL defines a collection of XML action elements that control the processing of the JSP document on the Web server. These elements have the same form as other elements of HTML and XML. One of the most commonly used JSTL control action elements is \texttt{if}, which specifies a Boolean expression as an attribute.\footnote{An attribute in HTML, which is embedded in the opening tag of an element, provides further information about that element.} The content of the \texttt{if} element (the text between the opening tag (<\texttt{if}>) and its closing tag (</\texttt{if}>)) is HTML code that will be included in the output document only if the Boolean expression evaluates to true. The \texttt{if} element is related to the C/C++ \#if preprocessor command. The JSP container processes the JSTL parts of JSP documents in a way that is similar to how the C/C++ preprocessor processes C and C++ programs. The preprocessor commands are instructions for the preprocessor to specify how the output file is to be constructed from the input file. Similarly, JSTL control action elements are instructions for the JSP processor to specify how to build the XML output file from the XML input file.

One common use of the \texttt{if} element is for the validation of form data submitted by a browser user. Form data is accessible by the JSP processor and can be tested with the \texttt{if} element to ensure that it is sensible data. If not, the \texttt{if} element can insert an error message for the user in the output document.

For multiple selection control, JSTL has \texttt{choose}, \texttt{when}, and \texttt{otherwise} elements. JSTL also includes a \texttt{forEach} element, which iterates over collections, which typically are form values from a client. The \texttt{forEach} element can include \texttt{begin}, \texttt{end}, and \texttt{step} attributes to control its iterations.

**Summary**

We have investigated the development of a number of programming languages. This chapter gives the reader a good perspective on current issues in language design. We have set the stage for an in-depth discussion of the important features of contemporary languages.

**Bibliographic Notes**

Perhaps the most important source of historical information about the development of early programming languages is *History of Programming Languages*, edited by Richard Wexelblat (1981). It contains the developmental background...
and environment of 13 important programming languages, as told by the designers themselves. A similar work resulted from a second “history” conference, published as a special issue of *ACM SIGPLAN Notices* (ACM, 1993a). In this work, the history and evolution of 13 more programming languages are discussed.

The paper “Early Development of Programming Languages” (Knuth and Pardo, 1977), which is part of the *Encyclopedia of Computer Science and Technology*, is an excellent 85-page work that details the development of languages up to and including Fortran. The paper includes example programs to demonstrate the features of many of those languages.

Another book of great interest is *Programming Languages: History and Fundamentals*, by Jean Sammet (1969). It is a 785-page work filled with details of 80 programming languages of the 1950s and 1960s. Sammet has also published several updates to her book, such as *Roster of Programming Languages for 1974–75* (1976).

### Review Questions

1. What is the meaning of Plankalkül? Who developed Plankalkül?
2. What was the simplest data type in Plankalkül?
3. How did Plankalkül implement iterations?
4. Speedcoding was invented to overcome two significant shortcomings of the computer hardware of the early 1950s. What were they?
5. Why was the technique used by Short Code called automatic programming?
6. In what year was IBM 704 introduced?
7. Where was Fortran developed? By whom?
8. Under what environmental constraints was Fortran developed?
9. How were data types specified for variables in Fortran I?
10. What was the most significant feature added to Fortran II to get Fortran IV?
11. What syntactic changes did Fortran 90 include?
12. Which version of Fortran was the first to support object-oriented programming?
13. Which version of Fortran was the first to have local scopes?
14. Why was Fortran list processing language (FLPL) designed and implemented as an extension to Fortran?
15. Where was Lisp developed? By whom?
16. In what ways do Scheme and metalanguage (ML) differ from each other?
17. What are the features of F# that make it unique?
18. What are the goals of the new language developed by GAMM and ACM?
19. In what version of ALGOL did the concept of data type appear?
20. What were the most important modifications made to ALGOL 58?
21. What change did ALGOL provide regarding input and output statements?
22. On what programming language was COBOL based?
23. In what year did the Basic design process begin?
24. How does COBOL allow variable names to be truly connotative?
25. What organization was most responsible for the early success of COBOL (in terms of extent of use)?
26. What were the objectives behind developing Basic?
27. Why is Alan Cooper known as “the father of Visual Basic”?
28. What facilities did PL/I provide?
29. What are the two fundamental characteristics shared by APL and SNOBOL?
30. What features of SIMULA 67 are now important parts of some object-oriented languages?
31. What innovation of data structuring was introduced in ALGOL 68 but is often credited to Pascal?
32. What was the primary deficiency of ALGOL 60?
33. Which language was used for teaching programming in the 1970s?
34. What operators in C were modeled on similar operators in ALGOL 68?
35. Why is C both liked and disliked?
36. What is a procedural language?
37. What are the two kinds of statements that populate a Prolog database?
38. What is the primary application area for which Ada was designed?
39. What are the concurrent program units of Ada called?
40. What are the different mechanisms in C that show that it was influenced by ALGOL 68?
41. Why did Ada 95 lose its popularity?
42. What is the meaning of operator overloading in C++?
43. What additional feature did the release 3.0 of C++ provide?
44. From what language does Objective-C borrow its syntax for method calls?
45. What is the primary application for Objective-C?
46. What do the Ada and COBOL languages have in common?
47. What are threads in Java?
48. What characteristic of Java is most evident in JavaScript?
49. How does the typing system of PHP and JavaScript differ from that of Java?
50. What array structure is included in C# but not in C, C++, or Java?
51. What are the characteristics of Perl's arrays?
52. For what application area is JavaScript most widely used?
53. What is the relationship between JavaScript and PHP, in terms of their use?
54. PHP's primary data structure is a combination of what two data structures from other languages?
55. What data structure does Python use in place of arrays?
56. What characteristic does Ruby share with Smalltalk?
57. What characteristic of Ruby's arithmetic operators makes them unique among those of other languages?
58. What is the similarity between Lua and JavaScript?
59. Is Lua normally compiled, purely interpreted, or impurely interpreted?
60. What deficiency of the `switch` statement of C is addressed with the changes made by C# to that statement?
61. What is the primary platform on which C# is used?
62. What are the inputs to an XSLT processor?
63. What is the output of an XSLT processor?
64. What element of the JSTL is related to a subprogram?
65. To what is a JSP document converted by a JSP processor?
66. Where are servlets executed?

**Problem Set**

1. What features of Plankalkül do you think would have had the greatest influence on Fortran 0 if the Fortran designers had been familiar with Plankalkül?
2. Determine the capabilities of Backus's 701 Speedcoding system, and compare them with those of a contemporary programmable hand calculator.
3. Write a short history of the A-0, A-1, and A-2 systems designed by Grace Hopper and her associates.
4. Compare the facilities of Fortran 0 with those of the Laning and Zierler system.
5. Which of the three original goals of the ALGOL design committee, in your opinion, was most difficult to achieve at that time?
6. Make an educated guess as to the most common syntax error in Lisp programs.
7. Lisp began as a pure functional language but gradually acquired more and more imperative features. Why?

8. Describe in detail the three most important reasons, in your opinion, why ALGOL 60 did not become a very widely used language.

9. Why, in your opinion, did COBOL allow long identifiers when Fortran and ALGOL did not?

10. Outline the major motivation of IBM in developing PL/I.

11. Was IBM’s assumption, on which it based its decision to develop PL/I, correct, given the history of computers and language developments since 1964?

12. Describe, in your own words, the concept of orthogonality in programming language design.

13. What is the primary reason why PL/I became more widely used than ALGOL 68?

14. What are the arguments both for and against the idea of a typeless language?

15. Are there any logic programming languages other than Prolog?

16. What is your opinion of the argument that languages that are too complex are too dangerous to use, and we should therefore keep all languages small and simple?

17. Do you think language design by committee is a good idea? Support your opinion.

18. Languages continually evolve. What sort of restrictions do you think are appropriate for changes in programming languages? Compare your answers with the evolution of Fortran.

19. Build a table identifying all of the major language developments, together with when they occurred, in what language they first appeared, and the identities of the developers.

20. There have been some public interchanges between Microsoft and Sun concerning the design of Microsoft’s J++ and C# and Sun’s Java. Read some of these documents, which are available on their respective Web sites, and write an analysis of the disagreements concerning the delegates.

21. In recent years data structures have evolved within scripting languages to replace traditional arrays. Explain the chronological sequence of these developments.

22. Explain two reasons why pure interpretation is an acceptable implementation method for several recent scripting languages.

23. Why, in your opinion, do new scripting languages appear more frequently than new compiled languages?

24. Give a brief general description of a markup-programming hybrid language.
1. To understand the value of records in a programming language, write a small program in a C-based language that uses an array of structs that store student information, including name, age, GPA as a float, and grade level as a string (e.g., “freshmen,” etc.). Also, write the same program in the same language without using structs.

2. To understand the value of recursion in a programming language, write a program that implements quicksort, first using recursion and then without recursion.

3. To understand the value of counting loops, write a program that implements matrix multiplication using counting loop constructs. Then write the same program using only logical loops—for example, while loops.
3.1 Introduction
3.2 The General Problem of Describing Syntax
3.3 Formal Methods of Describing Syntax
3.4 Attribute Grammars
3.5 Describing the Meanings of Programs: Dynamic Semantics
This chapter begins by defining the terms syntax and semantics. Then, a
detailed discussion of the most common method of describing syntax,
context-free grammars (also known as Backus-Naur Form), is presented.
Included in this discussion are derivations, parse trees, ambiguity, descriptions of
operator precedence and associativity, and extended Backus-Naur Form. Attribute
grammars, which can be used to describe both the syntax and static semantics of
programming languages, are discussed next. In the last section, three formal meth-
ods of describing semantics—operational, axiomatic, and denotational semantics—
are introduced. Because of the inherent complexity of the semantics description
methods, our discussion of them is brief. One could easily write an entire book on
just one of the three (as several authors have).

3.1 Introduction

The task of providing a concise yet understandable description of a program-
ning language is difficult but essential to the language’s success. ALGOL 60
and ALGOL 68 were first presented using concise formal descriptions; in both
cases, however, the descriptions were not easily understandable, partly because
each used a new notation. The levels of acceptance of both languages suffered
as a result. On the other hand, some languages have suffered the problem of
having many slightly different dialects, a result of a simple but informal and
imprecise definition.

One of the problems in describing a language is the diversity of the people
who must understand the description. Among these are initial evaluators,
implementors, and users. Most new programming languages are subjected to a
period of scrutiny by potential users, often people within the organization that
employs the language’s designer, before their designs are completed. These are
the initial evaluators. The success of this feedback cycle depends heavily on the
clarity of the description.

Programming language implementors obviously must be able to deter-
mine how the expressions, statements, and program units of a language are
formed, and also their intended effect when executed. The difficulty of the
implementors’ job is, in part, determined by the completeness and precision of
the language description.

Finally, language users must be able to determine how to encode software
solutions by referring to a language reference manual. Textbooks and courses
enter into this process, but language manuals are usually the only authoritative
printed information source about a language.

The study of programming languages, like the study of natural languages,
can be divided into examinations of syntax and semantics. The syntax of a
programming language is the form of its expressions, statements, and program
units. Its semantics is the meaning of those expressions, statements, and pro-
gram units. For example, the syntax of a Java while statement is

while (boolean_expr) statement
The semantics of this statement form is that when the current value of the Boolean expression is true, the embedded statement is executed. Then control implicitly returns to the Boolean expression to repeat the process. If the Boolean expression is false, control transfers to the statement following the while construct.

Although they are often separated for discussion purposes, syntax and semantics are closely related. In a well-designed programming language, semantics should follow directly from syntax; that is, the appearance of a statement should strongly suggest what the statement is meant to accomplish.

Describing syntax is easier than describing semantics, partly because a concise and universally accepted notation is available for syntax description, but none has yet been developed for semantics.

3.2 The General Problem of Describing Syntax

A language, whether natural (such as English) or artificial (such as Java), is a set of strings of characters from some alphabet. The strings of a language are called sentences or statements. The syntax rules of a language specify which strings of characters from the language's alphabet are in the language. English, for example, has a large and complex collection of rules for specifying the syntax of its sentences. By comparison, even the largest and most complex programming languages are syntactically very simple.

Formal descriptions of the syntax of programming languages, for simplicity's sake, often do not include descriptions of the lowest-level syntactic units. These small units are called lexemes. The description of lexemes can be given by a lexical specification, which is usually separate from the syntactic description of the language. The lexemes of a programming language include its numeric literals, operators, and special words, among others. One can think of programs as strings of lexemes rather than of characters.

Lexemes are partitioned into groups—for example, the names of variables, methods, classes, and so forth in a programming language form a group called identifiers. Each lexeme group is represented by a name, or token. So, a token of a language is a category of its lexemes. For example, an identifier is a token that can have lexemes, or instances, such as sum and total. In some cases, a token has only a single possible lexeme. For example, the token for the arithmetic operator symbol + has just one possible lexeme. Consider the following Java statement:

\[
\text{index} = 2 * \text{count} + 17;
\]

The lexemes and tokens of this statement are

<table>
<thead>
<tr>
<th>Lexemes</th>
<th>Tokens</th>
</tr>
</thead>
<tbody>
<tr>
<td>index</td>
<td>identifier</td>
</tr>
<tr>
<td>=</td>
<td>equal_sign</td>
</tr>
<tr>
<td>2</td>
<td>int Literal</td>
</tr>
</tbody>
</table>
Chapter 3  Describing Syntax and Semantics

*  mult_op
count  identifier
+  plus_op
17  int_literal
;  semicolon

The example language descriptions in this chapter are very simple, and most include lexeme descriptions.

3.2.1 Language Recognizers

In general, languages can be formally defined in two distinct ways: by recognition and by generation (although neither provides a definition that is practical by itself for people trying to learn or use a programming language). Suppose we have a language L that uses an alphabet Σ of characters. To define L formally using the recognition method, we would need to construct a mechanism R, called a recognition device, capable of reading strings of characters from the alphabet Σ. R would indicate whether a given input string was or was not in L. In effect, R would either accept or reject the given string. Such devices are like filters, separating legal sentences from those that are incorrectly formed. If R, when fed any string of characters over Σ, accepts it only if it is in L, then R is a description of L. Because most useful languages are, for all practical purposes, infinite, this might seem like a lengthy and ineffective process. Recognition devices, however, are not used to enumerate all of the sentences of a language—they have a different purpose.

The syntax analysis part of a compiler is a recognizer for the language the compiler translates. In this role, the recognizer need not test all possible strings of characters from some set to determine whether each is in the language. Rather, it need only determine whether given programs are in the language. In effect then, the syntax analyzer determines whether the given programs are syntactically correct. The structure of syntax analyzers, also known as parsers, is discussed in Chapter 4.

3.2.2 Language Generators

A language generator is a device that can be used to generate the sentences of a language. We can think of the generator as having a button that produces a sentence of the language every time it is pushed. Because the particular sentence that is produced by a generator when its button is pushed is unpredictable, a generator seems to be a device of limited usefulness as a language descriptor. However, people prefer certain forms of generators over recognizers because they can more easily read and understand them. By contrast, the syntax-checking portion of a compiler (a language recognizer) is not as useful a language description for a programmer because it can be used only in trial-and-error mode. For example, to determine the correct syntax of a particular statement using a compiler, the programmer can only submit a speculated version and note whether
the compiler accepts it. On the other hand, it is often possible to determine whether the syntax of a particular statement is correct by comparing it with the structure of the generator.

There is a close connection between formal generation and recognition devices for the same language. This was one of the seminal discoveries in computer science, and it led to much of what is now known about formal languages and compiler design theory. We return to the relationship of generators and recognizers in the next section.

### 3.3 Formal Methods of Describing Syntax

This section discusses the formal language-generation mechanisms, usually called grammars, that are commonly used to describe the syntax of programming languages.

#### 3.3.1 Backus-Naur Form and Context-Free Grammars

In the middle to late 1950s, two men, Noam Chomsky and John Backus, in unrelated research efforts, developed the same syntax description formalism, which subsequently became the most widely used method for programming language syntax.

##### 3.3.1.1 Context-Free Grammars

In the mid-1950s, Noam Chomsky, a noted linguist (among other things), described four classes of generative devices or grammars that define four classes of languages (Chomsky, 1956, 1959). Two of these grammar classes, named context-free and regular, turned out to be useful for describing the syntax of programming languages. The forms of the tokens of programming languages can be described by regular grammars. The syntax of whole programming languages, with minor exceptions, can be described by context-free grammars. Because Chomsky was a linguist, his primary interest was the theoretical nature of natural languages. He had no interest at the time in the artificial languages used to communicate with computers. So it was not until later that his work was applied to programming languages.

##### 3.3.1.2 Origins of Backus-Naur Form

Shortly after Chomsky’s work on language classes, the ACM-GAMM group began designing ALGOL 58. A landmark paper describing ALGOL 58 was presented by John Backus, a prominent member of the ACM-GAMM group, at an international conference in 1959 (Backus, 1959). This paper introduced a new formal notation for specifying programming language syntax. The new notation was later modified slightly by Peter Naur for the description of
ALGOL 60 (Naur, 1960). This revised method of syntax description became known as **Backus-Naur Form**, or simply **BNF**.

BNF is a natural notation for describing syntax. In fact, something similar to BNF was used by Panini to describe the syntax of Sanskrit several hundred years before Christ (Ingerman, 1967).

Although the use of BNF in the ALGOL 60 report was not immediately accepted by computer users, it soon became and is still the most popular method of concisely describing programming language syntax.

It is remarkable that BNF is nearly identical to Chomsky’s generative devices for context-free languages, called **context-free grammars**. In the remainder of the chapter, we refer to context-free grammars simply as grammars. Furthermore, the terms BNF and grammar are used interchangeably.

### 3.3.1.3 Fundamentals

A **metalanguage** is a language that is used to describe another language. BNF is a metalanguage for programming languages.

BNF uses abstractions for syntactic structures. A simple Java assignment statement, for example, might be represented by the abstraction `<assign>` (pointed brackets are often used to delimit names of abstractions). The actual definition of `<assign>` can be given by

\[ \langle \text{assign} \rangle \rightarrow \langle \text{var} \rangle = \langle \text{expression} \rangle \]

The text on the left side of the arrow, which is aptly called the **left-hand side** (LHS), is the abstraction being defined. The text to the right of the arrow is the definition of the LHS. It is called the **right-hand side** (RHS) and consists of some mixture of tokens, lexemes, and references to other abstractions. (Actually, tokens are also abstractions.) Altogether, the definition is called a **rule**, or **production**. In the example rule just given, the abstractions `<var>` and `<expression>` obviously must be defined for the `<assign>` definition to be useful.

This particular rule specifies that the abstraction `<assign>` is defined as an instance of the abstraction `<var>`, followed by the lexeme `=`, followed by an instance of the abstraction `<expression>`. One example sentence whose syntactic structure is described by the rule is

\[ \text{total} = \text{subtotal1} + \text{subtotal2} \]

The abstractions in a BNF description, or grammar, are often called **nonterminal symbols**, or simply **nonterminals**, and the lexemes and tokens of the rules are called **terminal symbols**, or simply **terminals**. A BNF description, or **grammar**, is a collection of rules.

Nonterminal symbols can have two or more distinct definitions, representing two or more possible syntactic forms in the language. Multiple definitions can be written as a single rule, with the different definitions separated
3.3 Formal Methods of Describing Syntax

by the symbol |, meaning logical OR. For example, a Java if statement can be described with the rules

\[
\begin{align*}
<\text{if_stmt}> & \rightarrow \text{if} \ ( <\text{logic_expr}> ) \ <\text{stmt}> \\
<\text{if_stmt}> & \rightarrow \text{if} \ ( <\text{logic_expr}> ) \ <\text{stmt}> \ \text{else} \ <\text{stmt}>
\end{align*}
\]

or with the rule

\[
<\text{if_stmt}> \rightarrow \text{if} \ ( <\text{logic_expr}> ) \ <\text{stmt}> \\
\quad \quad | \ \text{if} \ ( <\text{logic_expr}> ) \ <\text{stmt}> \ \text{else} \ <\text{stmt}>
\]

In these rules, <stmt> represents either a single statement or a compound statement.

Although BNF is simple, it is sufficiently powerful to describe nearly all of the syntax of programming languages. In particular, it can describe lists of similar constructs, the order in which different constructs must appear, and nested structures to any depth, and even imply operator precedence and operator associativity.

3.3.1.4 Describing Lists

Variable-length lists in mathematics are often written using an ellipsis (\ldots); 1, 2,\ldots is an example. BNF does not include the ellipsis, so an alternative method is required for describing lists of syntactic elements in programming languages (for example, a list of identifiers appearing on a data declaration statement). For BNF, the alternative is recursion. A rule is recursive if its LHS appears in its RHS. The following rules illustrate how recursion is used to describe lists:

\[
<\text{ident_list}> \rightarrow \text{identifier} \\
\quad \quad | \ \text{identifier, } <\text{ident_list}>
\]

This defines <ident_list> as either a single token (identifier) or an identifier followed by a comma and another instance of <ident_list>. Recursion is used to describe lists in many of the example grammars in the remainder of this chapter.

3.3.1.5 Grammars and Derivations

A grammar is a generative device for defining languages. The sentences of the language are generated through a sequence of applications of the rules, beginning with a special nonterminal of the grammar called the start symbol. This sequence of rule applications is called a derivation. In a grammar for a complete programming language, the start symbol represents a complete program and is often named <program>. The simple grammar shown in Example 3.1 is used to illustrate derivations.
The language described by the grammar of Example 3.1 has only one statement form: assignment. A program consists of the special word `begin`, followed by a list of statements separated by semicolons, followed by the special word `end`. An expression is either a single variable or two variables separated by either a `+` or `-` operator. The only variable names in this language are `A`, `B`, and `C`.

A derivation of a program in this language follows:

```
<program>    ->  begin stmt_list end
stmt_list    ->  stmt |
stmt         ->  var = expression |
var          ->  A | B | C |
expression   ->  var + var |
               ->  var - var |
               ->  var
```

This derivation, like all derivations, begins with the start symbol, in this case `<program>`. The symbol `=>` is read “derives.” Each successive string in the sequence is derived from the previous string by replacing one of the nonterminals with one of that nonterminal’s definitions. Each of the strings in the derivation, including `<program>`, is called a **sentential form**.

In this derivation, the replaced nonterminal is always the leftmost nonterminal in the previous sentential form. Derivations that use this order of replacement are called **leftmost derivations**. The derivation continues until the sentential form contains no nonterminals. That sentential form, consisting of only terminals, or lexemes, is the generated sentence.
In addition to leftmost, a derivation may be rightmost or in an order that is neither leftmost nor rightmost. Derivation order has no effect on the language generated by a grammar.

By choosing alternative RHSs of rules with which to replace nonterminals in the derivation, different sentences in the language can be generated. By exhaustively choosing all combinations of choices, the entire language can be generated. This language, like most others, is infinite, so one cannot generate all the sentences in the language in finite time.

Example 3.2 is another example of a grammar for part of a typical programming language.

### Example 3.2

**A Grammar for Simple Assignment Statements**

\[
\begin{align*}
\text{<assign>} & \rightarrow \text{id} = \text{expr} \\
\text{id} & \rightarrow A | B | C \\
\text{expr} & \rightarrow \text{id} + \text{expr} | \text{id} \times \text{expr} | ( \text{expr} ) | \text{id}
\end{align*}
\]

The grammar of Example 3.2 describes assignment statements whose right sides are arithmetic expressions with multiplication and addition operators and parentheses. For example, the statement

\[
A = B * ( A + C )
\]

is generated by the leftmost derivation:

\[
\begin{align*}
\text{<assign>} & \Rightarrow \text{id} = \text{expr} \\
& \Rightarrow A = \text{expr} \\
& \Rightarrow A = \text{id} * \text{expr} \\
& \Rightarrow A = B * \text{expr} \\
& \Rightarrow A = B * ( \text{expr} ) \\
& \Rightarrow A = B * ( \text{id} + \text{expr} ) \\
& \Rightarrow A = B * ( A + \text{expr} ) \\
& \Rightarrow A = B * ( A + \text{id} ) \\
& \Rightarrow A = B * ( A + C )
\end{align*}
\]

### 3.3.1.6 Parse Trees

One of the most attractive features of grammars is that they naturally describe the hierarchical syntactic structure of the sentences of the languages they define. These hierarchical structures are called **parse trees**. For example, the parse tree in Figure 3.1 shows the structure of the assignment statement derived previously.
Chapter 3 Describing Syntax and Semantics

Every internal node of a parse tree is labeled with a nonterminal symbol; every leaf is labeled with a terminal symbol. Every subtree of a parse tree describes one instance of an abstraction in the sentence.

3.3.1.7 Ambiguity

A grammar that generates a sentential form for which there are two or more distinct parse trees is said to be ambiguous. Consider the grammar shown in Example 3.3, which is a minor variation of the grammar shown in Example 3.2.

**Figure 3.1**
A parse tree for the simple statement
\[ A = B \ast (A + C) \]

---

**EXAMPLE 3.3** An Ambiguous Grammar for Simple Assignment Statements

\[
\begin{align*}
\langle \text{assign} \rangle & \rightarrow \langle \text{id} \rangle \ = \ \langle \text{expr} \rangle \\
\langle \text{id} \rangle & \rightarrow A \mid B \mid C \\
\langle \text{expr} \rangle & \rightarrow \langle \text{expr} \rangle \ + \ \langle \text{expr} \rangle \\
& \mid \langle \text{expr} \rangle \ * \ \langle \text{expr} \rangle \\
& \mid (\ \langle \text{expr} \rangle) \\
& \mid \langle \text{id} \rangle
\end{align*}
\]

The grammar of Example 3.3 is ambiguous because the sentence

\[ A = B + C \ast A \]

has two distinct parse trees, as shown in Figure 3.2. The ambiguity occurs because the grammar specifies slightly less syntactic structure than does the grammar of
Example 3.2. Rather than allowing the parse tree of an expression to grow only on the right, this grammar allows growth on both the left and the right.

Syntactic ambiguity of language structures is a problem because compilers often base the semantics of those structures on their syntactic form. Specifically, the compiler chooses the code to be generated for a statement by examining its parse tree. If a language structure has more than one parse tree, then the meaning of the structure cannot be determined uniquely. This problem is discussed in two specific examples in the following subsections.

There are several other characteristics of a grammar that are sometimes useful in determining whether a grammar is ambiguous. They include the following: (1) if the grammar generates a sentence with more than one leftmost derivation and (2) if the grammar generates a sentence with more than one rightmost derivation.

Some parsing algorithms can be based on ambiguous grammars. When such a parser encounters an ambiguous construct, it uses nongrammatical information provided by the designer to construct the correct parse tree. In many cases, an ambiguous grammar can be rewritten to be unambiguous but still generate the desired language.

### 3.3.1.8 Operator Precedence

When an expression includes two different operators, for example, $x + y * z$, one obvious semantic issue is the order of evaluation of the two operators (for example, in this expression is it add and then multiply, or vice versa?). This semantic question can be answered by assigning different precedence levels to operators. For example, if $*$ has been assigned higher precedence than $+$ (by the language designer), multiplication will be done first, regardless of the order of appearance of the two operators in the expression.

---

1. Note that it is mathematically impossible to determine whether an arbitrary grammar is ambiguous.
As stated previously, a grammar can describe a certain syntactic structure so that part of the meaning of the structure can be determined from its parse tree. In particular, the fact that an operator in an arithmetic expression is generated lower in the parse tree (and therefore must be evaluated first) can be used to indicate that it has precedence over an operator produced higher up in the tree. In the first parse tree of Figure 3.2, for example, the multiplication operator is generated lower in the tree, which could indicate that it has precedence over the addition operator in the expression. The second parse tree, however, indicates just the opposite. It appears, therefore, that the two parse trees indicate conflicting precedence information.

Notice that although the grammar of Example 3.2 is not ambiguous, the precedence order of its operators is not the usual one. In this grammar, a parse tree of a sentence with multiple operators, regardless of the particular operators involved, has the rightmost operator in the expression at the lowest point in the parse tree, with the other operators in the tree moving progressively higher as one moves to the left in the expression. For example, in the expression \( A + B \times C \), \( \times \) is the lowest in the tree, indicating it is to be done first. However, in the expression \( A \times B + C \), \( + \) is the lowest, indicating it is to be done first.

A grammar can be written for the simple expressions we have been discussing that is both unambiguous and specifies a consistent precedence of the \( + \) and \( \times \) operators, regardless of the order in which the operators appear in an expression. The correct ordering is specified by using separate nonterminal symbols to represent the operands of the operators that have different precedence. This requires additional nonterminals and some new rules. Instead of using <expr> for both operands of both \( + \) and \( \times \), we could use three nonterminals to represent operands, which allows the grammar to force different operators to different levels in the parse tree. If <expr> is the root symbol for expressions, \( + \) can be forced to the top of the parse tree by having <expr> directly generate only \( + \) operators, using the new nonterminal, <term>, as the right operand of \( + \). Next, we can define <term> to generate \( \times \) operators, using <term> as the left operand and a new nonterminal, <factor>, as its right operand. Now, \( \times \) will always be lower in the parse tree, simply because it is farther from the start symbol than \( + \) in every derivation. The grammar of Example 3.4 is such a grammar.

**Example 3.4** An Unambiguous Grammar for Expressions

\[
\begin{align*}
<assign> & \rightarrow <id> = <expr> \\
<id> & \rightarrow A | B | C \\
<expr> & \rightarrow <expr> + <term> \\
& \quad | <term> \\
<term> & \rightarrow <term> \times <factor> \\
& \quad | <factor> \\
<factor> & \rightarrow ( <expr> ) \\
& \quad | <id>
\end{align*}
\]
The grammar in Example 3.4 generates the same language as the grammars of Examples 3.2 and 3.3, but it is unambiguous and it specifies the usual precedence order of multiplication and addition operators. The following derivation of the sentence \( A = B + C \times A \) uses the grammar of Example 3.4:

\[
\begin{align*}
<\text{assign}> & \Rightarrow <\text{id}> = <\text{expr}> \\
& \Rightarrow A = <\text{expr}> \\
& \Rightarrow A = <\text{expr}> + <\text{term}> \\
& \Rightarrow A = <\text{term}> + <\text{term}> \\
& \Rightarrow A = <\text{factor}> + <\text{term}> \\
& \Rightarrow A = <\text{id}> + <\text{term}> \\
& \Rightarrow A = B + <\text{term}> \\
& \Rightarrow A = B + <\text{term}> \times <\text{factor}> \\
& \Rightarrow A = B + <\text{factor}> \times <\text{factor}> \\
& \Rightarrow A = B + <\text{id}> \times <\text{factor}> \\
& \Rightarrow A = B + C \times <\text{factor}> \\
& \Rightarrow A = B + C \times <\text{id}> \\
& \Rightarrow A = B + C \times A
\end{align*}
\]

The unique parse tree for this sentence, using the grammar of Example 3.4, is shown in Figure 3.3.

The connection between parse trees and derivations is very close: Either can easily be constructed from the other. Every derivation with an unambiguous grammar has a unique parse tree, although that tree can be represented by different derivations. For example, the following derivation of the sentence \( A = B + C \times A \) is different from the derivation of the same sentence given previously. This is a rightmost derivation, whereas the previous one is leftmost. Both of these derivations, however, are represented by the same parse tree.

\[
\begin{align*}
<\text{assign}> & \Rightarrow <\text{id}> = <\text{expr}> \\
& \Rightarrow <\text{id}> = <\text{expr}> + <\text{term}> \\
& \Rightarrow <\text{id}> = <\text{expr}> + <\text{term}> \times <\text{factor}> \\
& \Rightarrow <\text{id}> = <\text{expr}> + <\text{term}> \times <\text{id}> \\
& \Rightarrow <\text{id}> = <\text{expr}> + <\text{term}> \times A \\
& \Rightarrow <\text{id}> = <\text{expr}> + <\text{id}> \times A \\
& \Rightarrow <\text{id}> = <\text{expr}> + C \times A \\
& \Rightarrow <\text{id}> = <\text{term}> + C \times A \\
& \Rightarrow <\text{id}> = <\text{factor}> + C \times A \\
& \Rightarrow <\text{id}> = <\text{id}> + C \times A \\
& \Rightarrow <\text{id}> = B + C \times A \\
& \Rightarrow A = B + C \times A
\end{align*}
\]
3.3.1.9 Associativity of Operators

When an expression includes two operators that have the same precedence (as * and / usually have)—for example, \(A / B * C\)—a semantic rule is required to specify which should have precedence. This rule is named associativity. As was the case with precedence, a grammar for expressions may correctly imply operator associativity. Consider the following example of an assignment statement:

\[ A = B + C + A \]

The parse tree for this sentence, as defined with the grammar of Example 3.4, is shown in Figure 3.4.

The parse tree of Figure 3.4 shows the left addition operator lower than the right addition operator. This is the correct order if addition is meant to be left associative, which is typical. In most cases, the associativity of addition in a computer is irrelevant. In mathematics, addition is associative, which means that left and right associative orders of evaluation mean the same thing. That is, \((A + B) + C = A + (B + C)\). Floating-point addition in a computer, however, is not necessarily associative. For example, suppose floating-point values store seven digits of accuracy. Consider the problem of adding 11 numbers together, where one of the numbers is \(10^7\) and the other ten are 1. If the small numbers (the 1’s) are each added to the large number, one at a time, there is no effect on that number, because the small numbers occur in the eighth digit of the large number. However, if the small numbers are first added together and the result is

---

2. An expression with two occurrences of the same operator has the same issue; for example, \(A / B / C\).
Subtraction and division are not associative, whether in mathematics or in a computer. Therefore, correct associativity may be essential for an expression that contains either of them.

When a grammar rule has its LHS also appearing at the beginning of its RHS, the rule is said to be **left recursive**. This left recursion specifies left associativity. For example, the left recursion of the rules of the grammar of Example 3.4 causes it to make both addition and multiplication left associative. Unfortunately, left recursion disallows the use of some important syntax analysis algorithms. When one of these algorithms is to be used, the grammar must be modified to remove the left recursion. This, in turn, disallows the grammar from precisely specifying that certain operators are left associative. Fortunately, left associativity can be enforced by the compiler, even though the grammar does not dictate it.

In most languages that provide it, the exponentiation operator is right associative. To indicate right associativity, right recursion can be used. A grammar rule is **right recursive** if the LHS appears at the right end of the RHS. Rules such as

```
<factor> → <exp> ** <factor>
  | <exp>
<exp> → ( <expr> )
  | id
```

could be used to describe exponentiation as a right-associative operator.
### 3.3.1.10 An Unambiguous Grammar for *if-else*

The BNF rules for a Java *if-else* statement are as follows:

```
<if_stmt> → if (<logic_expr>) <stmt>
    if (<logic_expr>) <stmt> else <stmt>
```

If we also have `<stmt> → <if_stmt>`, this grammar is ambiguous. The simplest sentential form that illustrates this ambiguity is

```
if (<logic_expr>) if (<logic_expr>) <stmt> else <stmt>
```

The two parse trees in Figure 3.5 show the ambiguity of this sentential form. Consider the following example of this construct:

```java
if (done == true)
    if (denom == 0)
        quotient = 0;
    else quotient = num / denom;
```

The problem is that if the upper parse tree in Figure 3.5 is used as the basis for translation, the else clause would be executed when `done` is not true, which
probably is not what was intended by the author of the construct. We will examine the practical problems associated with this else-association problem in Chapter 8.

We will now develop an unambiguous grammar that describes this if statement. The rule for if constructs in many languages is that an else clause, when present, is matched with the nearest previous unmatched then clause. Therefore, there cannot be an if statement without an else between a then clause and its matching else. So, for this situation, statements must be distinguished between those that are matched and those that are unmatched, where unmatched statements are else-less ifs and all other statements are matched. The problem with the earlier grammar is that it treats all statements as if they had equal syntactic significance—that is, as if they were all matched.

To reflect the different categories of statements, different abstractions, or nonterminals, must be used. The unambiguous grammar based on these ideas follows:

\[
\begin{align*}
<\text{stmt}> & \rightarrow <\text{matched}> \mid <\text{unmatched}> \\
<\text{matched}> & \rightarrow \text{if} (<\text{logic}\_\text{expr}>) <\text{matched}> \text{else} <\text{matched}> \\
& \mid \text{any non-if statement} \\
<\text{unmatched}> & \rightarrow \text{if} (<\text{logic}\_\text{expr}>) <\text{stmt}> \\
& \mid \text{if} (<\text{logic}\_\text{expr}>) <\text{matched}> \text{else} <\text{unmatched}>
\end{align*}
\]

There is just one possible parse tree, using this grammar, for the following sentential form:

\[
\text{if} (<\text{logic}\_\text{expr}>) \text{if} (<\text{logic}\_\text{expr}>) <\text{stmt}> \text{else} <\text{stmt}>
\]

### 3.3.2 Extended BNF

Because of a few minor inconveniences in BNF, it has been extended in several ways. Most extended versions are called Extended BNF, or simply EBNF, even though they are not all exactly the same. The extensions do not enhance the descriptive power of BNF; they only increase its readability and writability.

Three extensions are commonly included in the various versions of EBNF. The first of these denotes an optional part of an RHS, which is delimited by brackets. For example, a C if-else statement can be described as

\[
<\text{if}\_\text{stmt}> \rightarrow \text{if} (<\text{expression}>) <\text{statement}> [\text{else} <\text{statement}>]
\]

Without the use of the brackets, the syntactic description of this statement would require the following two rules:

\[
<\text{if}\_\text{stmt}> \rightarrow \text{if} (<\text{expression}>) <\text{statement}> \\
\mid \text{if} (<\text{expression}>) <\text{statement}> \text{else} <\text{statement}>
\]

The second extension is the use of braces in an RHS to indicate that the enclosed part can be repeated indefinitely or left out altogether. This extension allows lists to be built with a single rule, instead of using recursion and two
rules. For example, lists of identifiers separated by commas can be described by the following rule:

\[ \text{ident_list} \rightarrow \text{identifier} \{, \text{identifier}\} \]

This is a replacement of the recursion by a form of implied iteration; the part enclosed within braces can be iterated any number of times.

The third common extension deals with multiple-choice options. When a single element must be chosen from a group, the options are placed in parentheses and separated by the OR operator, |. For example,

\[ \text{term} \rightarrow \text{term} (\times | / | \% ) \text{factor} \]

In BNF, a description of this \( \text{term} \) would require the following three rules:

\[ \begin{align*} 
\text{term} & \rightarrow \text{term} \ast \text{factor} \\
& | \text{term} / \text{factor} \\
& | \text{factor} 
\end{align*} \]

The brackets, braces, and parentheses in the EBNF extensions are meta-symbols, which means they are notational tools and not terminal symbols in the syntactic entities they help describe. In cases where these metasymbols are also terminal symbols in the language being described, the instances that are terminal symbols can be underlined or quoted. Example 3.5 illustrates the use of braces and multiple choices in an EBNF grammar.

**EXAMPLE 3.5** BNF and EBNF Versions of an Expression Grammar

**BNF:**

\[ \begin{align*} 
\text{expr} & \rightarrow \text{expr} + \text{term} \\
& | \text{expr} - \text{term} \\
& | \text{term} \\
\text{term} & \rightarrow \text{term} \ast \text{factor} \\
& | \text{term} / \text{factor} \\
& | \text{factor} \\
\text{factor} & \rightarrow \text{exp} ** \text{factor} \\
\text{exp} & \rightarrow (\text{expr}) \\
& | \text{id} 
\end{align*} \]

**EBNF:**

\[ \begin{align*} 
\text{expr} & \rightarrow \text{term} \{ (+ | - ) \text{term}\} \\
\text{term} & \rightarrow \text{factor} \{ (\ast | / ) \text{factor}\} \\
\text{factor} & \rightarrow \text{exp} \{ ** \text{exp}\} \\
\text{exp} & \rightarrow (\text{expr}) \\
& | \text{id} 
\end{align*} \]
3.3 Formal Methods of Describing Syntax

The BNF rule

\[ \text{<expr>} \rightarrow \text{<expr>} + \text{<term>} \]

clearly specifies—in fact forces—the + operator to be left associative. However, the EBNF version,

\[ \text{<expr>} \rightarrow \text{<term>} \{ + \text{<term>} \} \]

does not imply the direction of associativity. This problem is overcome in a syntax analyzer based on an EBNF grammar for expressions by designing the syntax analysis process to enforce the correct associativity. This is further discussed in Chapter 4.

Some versions of EBNF allow a numeric superscript to be attached to the right brace to indicate an upper limit to the number of times the enclosed part can be repeated. Also, some versions use a plus (+) superscript to indicate one or more repetitions. For example,

\[ \text{<compound>} \rightarrow \text{begin} \text{<stmt>} \{ \text{<stmt>} \} \text{end} \]

and

\[ \text{<compound>} \rightarrow \text{begin} \{ \text{<stmt>} \}^+ \text{end} \]

are equivalent.

In recent years, some variations on BNF and EBNF have appeared. Among these are the following:

- In place of the arrow, a colon is used and the RHS is placed on the next line.
- Instead of a vertical bar to separate alternative RHSs, they are simply placed on separate lines.
- In place of square brackets to indicate something being optional, the subscript opt is used. For example,

\[ \text{Constructor Declarator} \rightarrow \text{SimpleName} (\text{FormalParameterList}^{\text{opt}}) \]

- Rather than using the \( | \) symbol in a parenthesized list of elements to indicate a choice, the words “one of” are used. For example,

\[ \text{AssignmentOperator} \rightarrow \text{one of } = \ast = / \&= += -= \ll = \gg = \&= ^= \mid = \]

There is a standard for EBNF, ISO/IEC 14977:1996(1996), but it is rarely used. The standard uses the equal sign (=) instead of an arrow in rules, terminates each RHS with a semicolon, and requires quotes on all terminal symbols. It also specifies a host of other notational rules.

### 3.3.3 Grammars and Recognizers

Earlier in this chapter, we suggested that there is a close relationship between generation and recognition devices for a given language. In fact, given a context-free grammar, a recognizer for the language generated by the grammar can be
algorithmically constructed. A number of software systems have been developed that perform this construction. Such systems allow the quick creation of the syntax analysis part of a compiler for a new language and are therefore quite valuable. One of the first of these syntax analyzer generators is named yacc (yet another compiler compiler) (Johnson, 1975). There are now many such systems available.

### 3.4 Attribute Grammars

An attribute grammar is a device used to describe more of the structure of a programming language than can be described with a context-free grammar. An attribute grammar is an extension to a context-free grammar. The extension allows certain language rules to be conveniently described, such as type compatibility. Before we formally define the form of attribute grammars, we must clarify the concept of static semantics.

#### 3.4.1 Static Semantics

There are some characteristics of programming languages that are difficult to describe with BNF, and some that are impossible. As an example of a syntax rule that is difficult to specify with BNF, consider type compatibility rules. In Java, for example, a floating-point value cannot be assigned to an integer type variable, although the opposite is legal. Although this restriction can be specified in BNF, it requires additional nonterminal symbols and rules. If all of the typing rules of Java were specified in BNF, the grammar would become too large to be useful, because the size of the grammar determines the size of the syntax analyzer.

As an example of a syntax rule that cannot be specified in BNF, consider the common rule that all variables must be declared before they are referenced. It has been proven that this rule cannot be specified in BNF.

These problems exemplify the categories of language rules called static semantics rules. The static semantics of a language is only indirectly related to the meaning of programs during execution; rather, it has to do with the legal forms of programs (syntax rather than semantics). Many static semantic rules of a language state its type constraints. Static semantics is so named because the analysis required to check these specifications can be done at compile time.

Because of the problems of describing static semantics with BNF, a variety of more powerful mechanisms has been devised for that task. One such mechanism, attribute grammars, was designed by Knuth (1968a) to describe both the syntax and the static semantics of programs.

Attribute grammars are a formal approach both to describing and checking the correctness of the static semantics rules of a program. Although they are not
always used in a formal way in compiler design, the basic concepts of attribute grammars are at least informally used in every compiler (see Aho et al., 1986).

Dynamic semantics, which is the meaning of expressions, statements, and program units, is discussed in Section 3.5.

3.4.2 Basic Concepts

Attribute grammars are context-free grammars to which have been added attributes, attribute computation functions, and predicate functions. Attributes, which are associated with grammar symbols (the terminal and nonterminal symbols), are similar to variables in the sense that they can have values assigned to them. Attribute computation functions, sometimes called semantic functions, are associated with grammar rules. They are used to specify how attribute values are computed. Predicate functions, which state the static semantic rules of the language, are associated with grammar rules.

These concepts will become clearer after we formally define attribute grammars and provide an example.

3.4.3 Attribute Grammars Defined

An attribute grammar is a grammar with the following additional features:

- Associated with each grammar symbol X is a set of attributes A(X). The set A(X) consists of two disjoint sets S(X) and I(X), called synthesized and inherited attributes, respectively. Synthesized attributes are used to pass semantic information up a parse tree, while inherited attributes pass semantic information down and across a tree.

- Associated with each grammar rule is a set of semantic functions and a possibly empty set of predicate functions over the attributes of the symbols in the grammar rule. For a rule \( X_0 \rightarrow X_1 \ldots X_n \), the synthesized attributes of \( X_0 \) are computed with semantic functions of the form \( S(X_0) = f(A(X_1), \ldots, A(X_n)) \). So the value of a synthesized attribute on a parse tree node depends only on the values of the attributes on that node’s children nodes. Inherited attributes of symbols \( X_j, 1 \leq j \leq n \) (in the rule above), are computed with a semantic function of the form \( I(X_j) = f(A(X_0), \ldots, A(X_{j-1})) \). So the value of an inherited attribute on a parse tree node depends on the attribute values of that node’s parent node and those of its sibling nodes. Note that, to avoid circularity, inherited attributes are often restricted to functions of the form \( I(X_j) = f(A(X_0), \ldots, A(X_{j-1})) \). This form prevents an inherited attribute from depending on itself or on attributes to the right in the parse tree.

- A predicate function has the form of a Boolean expression on the union of the attribute set \( \{A(X_0), \ldots, A(X_n)\} \) and a set of literal attribute values. The only derivations allowed with an attribute grammar are those in which every predicate associated with every nonterminal is true. A false predicate function value indicates a violation of the syntax or static semantics rules of the language.
A parse tree of an attribute grammar is the parse tree based on its underlying BNF grammar, with a possibly empty set of attribute values attached to each node. If all the attribute values in a parse tree have been computed, the tree is said to be fully attributed. Although in practice it is not always done this way, it is convenient to think of attribute values as being computed after the complete unattributed parse tree has been constructed by the compiler.

### 3.4.4 Intrinsic Attributes

**Intrinsic attributes** are synthesized attributes of leaf nodes whose values are determined outside the parse tree. For example, the type of an instance of a variable in a program could come from the symbol table, which is used to store variable names and their types. The contents of the symbol table are set based on earlier declaration statements. Initially, assuming that an unattributed parse tree has been constructed and that attribute values are needed, the only attributes with values are the intrinsic attributes of the leaf nodes. Given the intrinsic attribute values on a parse tree, the semantic functions can be used to compute the remaining attribute values.

### 3.4.5 Examples of Attribute Grammars

As a very simple example of how attribute grammars can be used to describe static semantics, consider the following fragment of an attribute grammar that describes the rule that the name on the `end` of an Ada procedure must match the procedure’s name. (This rule cannot be stated in BNF.) The string attribute of `<proc_name>`, denoted by `<proc_name>.string`, is the actual string of characters that were found immediately following the reserved word `procedure` by the compiler. Notice that when there is more than one occurrence of a nonterminal in a syntax rule in an attribute grammar, the nonterminals are subscripted with brackets to distinguish them. Neither the subscripts nor the brackets are part of the described language.

**Syntax rule:**

\[
<\text{proc_def}> \rightarrow \text{procedure} <\text{proc_name}> [1] \\
<\text{proc_body}> \text{end} <\text{proc_name}> [2] ;
\]

**Predicate:**

\[
<\text{proc_name}> [1] \text{string} == <\text{proc_name}> [2] .\text{string}
\]

In this example, the predicate rule states that the name string attribute of the `<proc_name>` nonterminal in the subprogram header must match the name string attribute of the `<proc_name>` nonterminal following the end of the subprogram.

Next, we consider a larger example of an attribute grammar. In this case, the example illustrates how an attribute grammar can be used to check the type rules of a simple assignment statement. The syntax and static semantics of this assignment statement are as follows: The only variable names are `A`, `B`, and `C`. The right side of the assignments can be either a variable or an expression in the form of a variable added to another variable. The variables can be one of two types: int or real. When there are two variables on the right side of an assignment, they need not be the same type. The type of the expression

\[
\text{Syntax rule: } <\text{proc_def}> \rightarrow <\text{proc_name}> [1] \\
<\text{proc_body}> \text{end} <\text{proc_name}> [2] ;
\]

**Predicate:**

\[
<\text{proc_name}> [1] \text{string} == <\text{proc_name}> [2] .\text{string}
\]
when the operand types are not the same is always real. When they are the same, the expression type is that of the operands. The type of the left side of the assignment must match the type of the right side. So the types of operands in the right side can be mixed, but the assignment is valid only if the target and the value resulting from evaluating the right side have the same type. The attribute grammar specifies these static semantic rules.

The syntax portion of our example attribute grammar is

\[
\begin{align*}
\text{<assign>} & \rightarrow \text{<var>} = \text{<expr> } \\
\text{<expr>} & \rightarrow \text{<var>} + \text{<var>} \\ & \quad | \quad \text{<var>} \\
\text{<var>} & \rightarrow A \mid B \mid C
\end{align*}
\]

The attributes for the nonterminals in the example attribute grammar are described in the following paragraphs:

- **actual_type**—A synthesized attribute associated with the nonterminals `<var>` and `<expr>`. It is used to store the actual type, int or real, of a variable or expression. In the case of a variable, the actual type is intrinsic. In the case of an expression, it is determined from the actual types of the child node or children nodes of the `<expr>` nonterminal.

- **expected_type**—An inherited attribute associated with the nonterminal `<expr>`. It is used to store the type, either int or real, that is expected for the expression, as determined by the type of the variable on the left side of the assignment statement.

The complete attribute grammar follows in Example 3.6.

---

**Example 3.6**

**An Attribute Grammar for Simple Assignment Statements**

1. Syntax rule: `<assign> → <var> = <expr>`
   Semantic rule: `<expr>.expected_type ← <var>.actual_type`
   Semantic rule: `<expr>.actual_type ←`
   ```
   \begin{align*}
   & \text{if } (<var>[2].actual_type = \text{int}) \text{ and} \\
   & \quad (<var>[3].actual_type = \text{int}) \\
   & \text{then int} \\
   & \text{else real} \\
   & \text{end if}
   \end{align*}
   ```
   Predicate: `<expr>.actual_type == <expr>.expected_type`
3. Syntax rule: `<expr> → <var>`
   Semantic rule: `<expr>.actual_type ← <var>.actual_type`
   Predicate: `<expr>.actual_type == <expr>.expected_type`
4. Syntax rule: `<var> → A | B | C`
   Semantic rule: `<var>.actual_type ← \text{look-up (<var>.string)}`

The look-up function looks up a given variable name in the symbol table and returns the variable’s type.
A parse tree of the sentence \( A = A + B \) generated by the grammar in Example 3.6 is shown in Figure 3.6. As in the grammar, bracketed numbers are added after the repeated node labels in the tree so they can be referenced unambiguously.

### 3.4.6 Computing Attribute Values

Now, consider the process of computing the attribute values of a parse tree, which is sometimes called **decorating** the parse tree. If all attributes were inherited, this could proceed in a completely top-down order, from the root to the leaves. Alternatively, it could proceed in a completely bottom-up order, from the leaves to the root, if all the attributes were synthesized. Because our grammar has both synthesized and inherited attributes, the evaluation process cannot be in any single direction. The following is an evaluation of the attributes, in an order in which it is possible to compute them:

1. \(<\text{var}>.\text{actual_type} \leftarrow \text{look-up}(A)\) (Rule 4)
2. \(<\text{expr}>.\text{expected_type} \leftarrow <\text{var}>.\text{actual_type}\) (Rule 1)
3. \(<\text{var}[2]>.\text{actual_type} \leftarrow \text{look-up}(A)\) (Rule 4)
   \(<\text{var}[3]>.\text{actual_type} \leftarrow \text{look-up}(B)\) (Rule 4)
4. \(<\text{expr}>.\text{actual_type} \leftarrow \text{either int or real}\) (Rule 2)
5. \(<\text{expr}>.\text{expected_type} == <\text{expr}>.\text{actual_type}\) is either
   TRUE or FALSE (Rule 2)

The tree in Figure 3.7 shows the flow of attribute values in the example of Figure 3.6. Solid lines show the parse tree; dashed lines show attribute flow in the tree.

The tree in Figure 3.8 shows the final attribute values on the nodes. In this example, \( A \) is defined as a real and \( B \) is defined as an int.

Determining attribute evaluation order for the general case of an attribute grammar is a complex problem, requiring the construction of a dependency graph to show all attribute dependencies.

### Figure 3.6

A parse tree for
\[ A = A + B \]
3.4.7 Evaluation

Checking the static semantic rules of a language is an essential part of all compilers. Even if a compiler writer has never heard of an attribute grammar, he or she would need to use their fundamental ideas to design the checks of static semantics rules for his or her compiler.

One of the main difficulties in using an attribute grammar to describe all of the syntax and static semantics of a real contemporary programming language is the size and complexity of the attribute grammar. The large number of attributes and semantic rules required for a complete programming language make such grammars difficult to write and read. Furthermore, the attribute values on a large parse tree are costly to evaluate. On the other hand, less formal attribute
grammars are a powerful and commonly used tool for compiler writers, who are more interested in the process of producing a compiler than they are in formalism.

### 3.5 Describing the Meanings of Programs: Dynamic Semantics

We now turn to the difficult task of describing the **dynamic semantics**, or meaning, of the expressions, statements, and program units of a programming language. Because of the power and naturalness of the available notation, describing syntax is a relatively simple matter. On the other hand, no universally accepted notation or approach has been devised for dynamic semantics. In this section, we briefly describe several of the methods that have been developed. For the remainder of this section, when we use the term *semantics*, we mean dynamic semantics.

There are several different reasons underlying the need for a methodology and notation for describing semantics. Programmers obviously need to know precisely what the statements of a language do before they can use them effectively in their programs. Compiler writers must know exactly what language constructs mean to design implementations for them correctly. If there were a precise semantics specification of a programming language, programs written in the language potentially could be proven correct without testing. Also, compilers could be shown to produce programs that exhibited exactly the behavior given in the language definition; that is, their correctness could be verified. A complete specification of the syntax and semantics of a programming language could be used by a tool to generate a compiler for the language automatically. Finally, language designers, who would develop the semantic descriptions of their languages, could in the process discover ambiguities and inconsistencies in their designs.

Software developers and compiler designers typically determine the semantics of programming languages by reading English explanations in language manuals. Because such explanations are often imprecise and incomplete, this approach is clearly unsatisfactory. Due to the lack of complete semantics specifications of programming languages, programs are rarely proven correct without testing, and commercial compilers are never generated automatically from language descriptions.

Scheme, a functional language described in Chapter 15, is one of only a few programming languages whose definition includes a formal semantics description. However, the method used is not one described in this chapter, as this chapter is focused on approaches that are suitable for imperative languages.

#### 3.5.1 Operational Semantics

The idea behind **operational semantics** is to describe the meaning of a statement or program by specifying the effects of running it on a machine. The effects on the machine are viewed as the sequence of changes in its state, where
the machine’s state is the collection of the values in its storage. An obvious operational semantics description, then, is given by executing a compiled version of the program on a computer. Most programmers have, on at least one occasion, written a small test program to determine the meaning of some programming language construct, often while learning the language. Essentially, what such a programmer is doing is using operational semantics to determine the meaning of the construct.

There are several problems with using this approach for complete formal semantics descriptions. First, the individual steps in the execution of machine language and the resulting changes to the state of the machine are too small and too numerous. Second, the storage of a real computer is too large and complex. There are usually several levels of memory devices, as well as connections to enumerable other computers and memory devices through networks. Therefore, machine languages and real computers are not used for formal operational semantics. Rather, intermediate-level languages and interpreters for idealized computers are designed specifically for the process.

There are different levels of uses of operational semantics. At the highest level, the interest is in the final result of the execution of a complete program. This is sometimes called natural operational semantics. At the lowest level, operational semantics can be used to determine the precise meaning of a program through an examination of the complete sequence of state changes that occur when the program is executed. This use is sometimes called structural operational semantics.

3.5.1.1 The Basic Process

The first step in creating an operational semantics description of a language is to design an appropriate intermediate language, where the primary desired characteristic of the language is clarity. Every construct of the intermediate language must have an obvious and unambiguous meaning. This language is at the intermediate level, because machine language is too low-level to be easily understood and another high-level language is obviously not suitable. If the semantics description is to be used for natural operational semantics, a virtual machine (an interpreter) must be constructed for the intermediate language. The virtual machine can be used to execute either single statements, code segments, or whole programs. The semantics description can be used without a virtual machine if the meaning of a single statement is all that is required. In this use, which is structural operational semantics, the intermediate code can be visually inspected.

The basic process of operational semantics is not unusual. In fact, the concept is frequently used in programming textbooks and programming language reference manuals. For example, the semantics of the C `for` construct can be described in terms of simpler statements, as in
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C Statement
for (expr1; expr2; expr3) {
  ...
}

Meaning
expr1;
loop: if expr2 == 0 goto out
  ...
expr3;
  goto loop
out: ...

The human reader of such a description is the virtual computer and is assumed to be able to “execute” the instructions in the definition correctly and recognize the effects of the “execution.”

The intermediate language and its associated virtual machine used for formal operational semantics descriptions are often highly abstract. The intermediate language is meant to be convenient for the virtual machine, rather than for human readers. For our purposes, however, a more human-oriented intermediate language could be used. As such an example, consider the following list of statements, which would be adequate for describing the semantics of the simple control statements of a typical programming language:

ident = var
ident = ident + 1
ident = ident – 1
goto label
if var relop var goto label

In these statements, relop is one of the relational operators from the set {=, <>, >, <, >=, <=}, ident is an identifier, and var is either an identifier or a constant. These statements are all simple and therefore easy to understand and implement.

A slight generalization of these three assignment statements allows more general arithmetic expressions and assignment statements to be described. The new statements are

ident = var bin_op var
ident = un_op var

where bin_op is a binary arithmetic operator and un_op is a unary operator. Multiple arithmetic data types and automatic type conversions, of course, complicate this generalization. Adding just a few more relatively simple instructions would allow the semantics of arrays, records, pointers, and subprograms to be described.

In Chapter 8, the semantics of various control statements are described using this intermediate language.

3.5.1.2 Evaluation

The first and most significant use of formal operational semantics was to describe the semantics of PL/I (Wegner, 1972). That particular abstract machine and the translation rules for PL/I were together named the Vienna Definition Language (VDL), after the city where IBM designed it.
Operational semantics provides an effective means of describing semantics for language users and language implementors, as long as the descriptions are kept simple and informal. The VDL description of PL/I, unfortunately, is so complex that it serves no practical purpose.

Operational semantics depends on programming languages of lower levels, not mathematics. The statements of one programming language are described in terms of the statements of a lower-level programming language. This approach can lead to circularities, in which concepts are indirectly defined in terms of themselves. The methods described in the following two sections are much more formal, in the sense that they are based on mathematics and logic, not programming languages.

3.5.2 Denotational Semantics

Denotational semantics is the most rigorous and most widely known formal method for describing the meaning of programs. It is solidly based on recursive function theory. A thorough discussion of the use of denotational semantics to describe the semantics of programming languages is necessarily long and complex. It is our intent to provide the reader with an introduction to the central concepts of denotational semantics, along with a few simple examples that are relevant to programming language specifications.

The process of constructing a denotational semantics specification for a programming language requires one to define for each language entity both a mathematical object and a function that maps instances of that language entity onto instances of the mathematical object. Because the objects are rigorously defined, they model the exact meaning of their corresponding entities. The idea is based on the fact that there are rigorous ways of manipulating mathematical objects but not programming language constructs. The difficulty with this method lies in creating the objects and the mapping functions. The method is named denotational because the mathematical objects denote the meaning of their corresponding syntactic entities.

The mapping functions of a denotational semantics programming language specification, like all functions in mathematics, have a domain and a range. The domain is the collection of values that are legitimate parameters to the function; the range is the collection of objects to which the parameters are mapped. In denotational semantics, the domain is called the syntactic domain, because it is syntactic structures that are mapped. The range is called the semantic domain.

Denotational semantics is related to operational semantics. In operational semantics, programming language constructs are translated into simpler programming language constructs, which become the basis of the meaning of the construct. In denotational semantics, programming language constructs are mapped to mathematical objects, either sets or, more often, functions. However, unlike operational semantics, denotational semantics does not model the step-by-step computational processing of programs.
3.5.2.1 Two Simple Examples

We use a very simple language construct, character string representations of binary numbers, to introduce the denotational method. The syntax of such binary numbers can be described by the following grammar rules:

\[\begin{align*}
<\text{bin\_num}> & \rightarrow '0' \\
& \mid '1' \\
& \mid <\text{bin\_num}> '0' \\
& \mid <\text{bin\_num}> '1'
\end{align*}\]

A parse tree for the example binary number, 110, is shown in Figure 3.9. Notice that we put apostrophes around the syntactic digits to show they are not mathematical digits. This is similar to the relationship between ASCII coded digits and mathematical digits. When a program reads a number as a string, it must be converted to a mathematical number before it can be used as a value in the program.

![Figure 3.9](image)

The syntactic domain of the mapping function for binary numbers is the set of all character string representations of binary numbers. The semantic domain is the set of nonnegative decimal numbers, symbolized by \(N\).

To describe the meaning of binary numbers using denotational semantics, we associate the actual meaning (a decimal number) with each rule that has a single terminal symbol as its RHS.

In our example, decimal numbers must be associated with the first two grammar rules. The other two grammar rules are, in a sense, computational rules, because they combine a terminal symbol, to which an object can be associated, with a nonterminal, which can be expected to represent some construct. Presuming an evaluation that progresses upward in the parse tree, the nonterminal in the right side would already have its meaning attached. So, a syntax rule with a nonterminal as its RHS would require a function that computed the meaning of the LHS, which represents the meaning of the complete RHS.

The semantic function, named \(M_{\text{bin}}\), maps the syntactic objects, as described in the previous grammar rules, to the objects in \(N\), the set of nonnegative decimal numbers. The function \(M_{\text{bin}}\) is defined as follows:
In part because we need it later, we now show a similar example for describing the meaning of syntactic decimal literals. In this case, the syntactic domain is the set of character string representations of decimal numbers. The semantic domain is once again the set $\mathbb{N}$.

$$<\text{dec_num}> \rightarrow '0' | '1' | '2' | '3' | '4' | '5' | '6' | '7' | '8' | '9' | <\text{dec_num}> ('0' | '1' | '2' | '3' | '4' | '5' | '6' | '7' | '8' | '9')$$

The denotational mappings for these syntax rules are

- $M_{\text{dec}}('0') = 0$
- $M_{\text{dec}}('1') = 1$
- $M_{\text{dec}}(<\text{dec_num}> '0') = 10 \times M_{\text{dec}}(<\text{dec_num}>)$
- $M_{\text{dec}}(<\text{dec_num}> '1') = 10 \times M_{\text{dec}}(<\text{dec_num}>) + 1$
- $\ldots$
- $M_{\text{dec}}(<\text{dec_num}> '9') = 10 \times M_{\text{dec}}(<\text{dec_num}>) + 9$

In the following sections, we present the denotational semantics descriptions of a few simple constructs. The most important simplifying assumption made here is that both the syntax and static semantics of the constructs are correct. In addition, we assume that only two scalar types are included: integer and Boolean.
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3.5.2.2 The State of a Program

The denotational semantics of a program could be defined in terms of state changes in an ideal computer. Operational semantics are defined in this way, and denotational semantics are defined in nearly the same way. In a further simplification, however, denotational semantics is defined in terms of only the values of all of the program’s variables. So, denotational semantics uses the state of the program to describe meaning, whereas operational semantics uses the state of a machine. The key difference between operational semantics and denotational semantics is that state changes in operational semantics are defined by coded algorithms, written in some programming language, whereas in denotational semantics, state changes are defined by mathematical functions.

Let the state $s$ of a program be represented as a set of ordered pairs, as follows:

$$s = \{ <i_1, v_1>, <i_2, v_2>, \ldots, <i_n, v_n> \}$$

Each $i$ is the name of a variable, and the associated $v$'s are the current values of those variables. Any of the $v$'s can have the special value $\text{undef}$, which indicates that its associated variable is currently undefined. Let $\text{VARMAP}$ be a function of two parameters: a variable name and the program state. The value of $\text{VARMAP} (i_j, s)$ is $v_j$ (the value paired with $i_j$ in state $s$). Most semantics mapping functions for programs and program constructs map states to states. These state changes are used to define the meanings of programs and program constructs. Some language constructs—for example, expressions—are mapped to values, not states.

3.5.2.3 Expressions

Expressions are fundamental to most programming languages. We assume here that expressions have no side effects. Furthermore, we deal with only very simple expressions: The only operators are $+$ and $\ast$, and an expression can have at most one operator; the only operands are scalar integer variables and integer literals; there are no parentheses; and the value of an expression is an integer.

Following is the BNF description of these expressions:

$$<\text{expr}> \rightarrow <\text{dec}\_\text{num}> | <\text{var}> | <\text{binary}\_\text{expr}>$$  
$$<\text{binary}\_\text{expr}> \rightarrow <\text{left}\_\text{expr}> <\text{operator}> <\text{right}\_\text{expr}>$$  
$$<\text{left}\_\text{expr}> \rightarrow <\text{dec}\_\text{num}> | <\text{var}>$$  
$$<\text{right}\_\text{expr}> \rightarrow <\text{dec}\_\text{num}> | <\text{var}>$$  
$$<\text{operator}> \rightarrow + | *$$

The only error we consider in expressions is a variable having an undefined value. Obviously, other errors can occur, but most of them are machine-dependent. Let $\mathbb{Z}$ be the set of integers, and let $\text{error}$ be the error value. Then $\mathbb{Z} \cup \{\text{error}\}$ is the semantic domain for the denotational specification for our expressions.
The mapping function for a given expression E and state s follows. To distinguish between mathematical function definitions and the assignment statements of programming languages, we use the symbol $\Delta$ to define mathematical functions. The implication symbol, $=>$, used in this definition connects the form of an operand with its associated case (or switch) construct. Dot notation is used to refer to the child nodes of a node. For example, $\langle\text{binary Expr}.\langle\text{left Expr}\rangle\rangle$ refers to the left child node of $\langle\text{binary Expr}\rangle$.

$M_e (\langle\text{Expr}\rangle, s) \Delta = \text{case } \langle\text{Expr}\rangle \text{ of}$

- $\langle\text{dec num}\rangle \Rightarrow M_{\text{dec}} (\langle\text{dec num}\rangle, s)$
- $\langle\text{var}\rangle \Rightarrow \text{if } \text{VARMAP (\langle var\rangle, s) } == \text{undef}$
  - then error
  - else $\text{VARMAP (\langle var\rangle, s)}$
- $\langle\text{binary Expr}\rangle \Rightarrow$
  - if ($M_e (\langle\text{binary Expr}.\langle\text{left Expr}\rangle,s) \text{ == undef OR } M_e (\langle\text{binary Expr}.\langle\text{right Expr}\rangle,s) \text{ == undef}$)
    - then error
  - else if ($\langle\text{binary Expr}.\langle\text{operator} == '+'\rangle\rangle$
    - then $M_e (\langle\text{binary Expr}.\langle\text{left Expr}\rangle,s) +$
      - $M_e (\langle\text{binary Expr}.\langle\text{right Expr}\rangle,s)$
    - else $M_e (\langle\text{binary Expr}.\langle\text{left Expr}\rangle,s) \times$
      - $M_e (\langle\text{binary Expr}.\langle\text{right Expr}\rangle,s)$

### 3.5.2.4 Assignment Statements

An assignment statement is an expression evaluation plus the setting of the target variable to the expression’s value. In this case, the meaning function maps a state to a state. This function can be described with the following:

$M_a (x = E, s) \Delta = \text{if } M_e (E, s) == \text{error}$

- then error
- else $s' = \{<i_1, v_1'>, <i_2, v_2'>, \ldots, <i_n, v_n'>\}$, where
  - for $j = 1, 2, \ldots, n$
    - if $i_j == x$
      - then $v_j' = M_e (E, s)$
    - else $v_j' = \text{VARMAP (i_j, s)}$

Note that the comparison in the third last line above, $i_j == x$, is of names, not values.

### 3.5.2.5 Logical Pretest Loops

The denotational semantics of a logical pretest loop is deceptively simple. To expedite the discussion, we assume that there are two other existing mapping functions, $M_{\text{sl}}$ and $M_{\text{bn}}$, that map statement lists and states to states and Boolean expressions to Boolean values (or error), respectively. The function is
\[ M_1(\text{while } B \text{ do } L, s) \Delta= \text{if } M_b(B, s) == \text{undef} \]
\[ \text{then } \text{error} \]
\[ \text{else if } M_b(B, s) == \text{false} \]
\[ \text{then } s \]
\[ \text{else if } M_{sl}(L, s) == \text{error} \]
\[ \text{then } \text{error} \]
\[ \text{else } M_1(\text{while } B \text{ do } L, M_{sl}(L, s)) \]

The meaning of the loop is simply the value of the program variables after the statements in the loop have been executed the prescribed number of times, assuming there have been no errors. In essence, the loop has been converted from iteration to recursion, where the recursion control is mathematically defined by other recursive state mapping functions. Recursion is easier to describe with mathematical rigor than iteration.

One significant observation at this point is that this definition, like actual program loops, may compute nothing because of nontermination.

### 3.5.2.6 Evaluation

Objects and functions, such as those used in the earlier constructs, can be defined for the other syntactic entities of programming languages. When a complete system has been defined for a given language, it can be used to determine the meaning of complete programs in that language. This provides a framework for thinking about programming in a highly rigorous way.

As stated previously, denotational semantics can be used as an aid to language design. For example, statements for which the denotational semantic description is complex and difficult may indicate to the designer that such statements may also be difficult for language users to understand and that an alternative design may be in order.

Because of the complexity of denotational descriptions, they are of little use to language users. On the other hand, they provide an excellent way to describe a language concisely.

Although the use of denotational semantics is normally attributed to Scott and Strachey (1971), the general denotational approach to language description can be traced to the nineteenth century (Frege, 1892).

### 3.5.3 Axiomatic Semantics

Axiomatic semantics, thus named because it is based on mathematical logic, is the most abstract approach to semantics specification discussed in this chapter. Rather than directly specifying the meaning of a program, axiomatic semantics specifies what can be proven about the program. Recall that one of the possible uses of semantic specifications is to prove the correctness of programs.
In axiomatic semantics, there is no model of the state of a machine or program or model of state changes that take place when the program is executed. The meaning of a program is based on relationships among program variables and constants, which are the same for every execution of the program.

Axiomatic semantics has two distinct applications: program verification and program semantics specification. This section focuses on program verification in its description of axiomatic semantics.

Axiomatic semantics was defined in conjunction with the development of an approach to proving the correctness of programs. Such correctness proofs, when they can be constructed, show that a program performs the computation described by its specification. In a proof, each statement of a program is both preceded and followed by a logical expression that specifies constraints on program variables. These, rather than the entire state of an abstract machine (as with operational semantics), are used to specify the meaning of the statement. The notation used to describe constraints—indeed, the language of axiomatic semantics—is predicate calculus. Although simple Boolean expressions are often adequate to express constraints, in some cases they are not.

When axiomatic semantics is used to specify formally the meaning of a statement, the meaning is defined by the statement’s effect on assertions about the data affected by the statement.

### 3.5.3.1 Assertions

The logical expressions used in axiomatic semantics are called predicates, or assertions. An assertion immediately preceding a program statement describes the constraints on the program variables at that point in the program. An assertion immediately following a statement describes the new constraints on those variables (and possibly others) after execution of the statement. These assertions are called the **precondition** and **postcondition**, respectively, of the statement. For two adjacent statements, the postcondition of the first serves as the precondition of the second. Developing an axiomatic description or proof of a given program requires that every statement in the program has both a precondition and a postcondition.

In the following sections, we examine assertions from the point of view that preconditions for statements are computed from given postconditions, although it is possible to consider these in the opposite sense. We assume all variables are integer type. As a simple example, consider the following assignment statement and postcondition:

\[
\text{sum} = 2 * \text{x} + 1 \quad (\text{sum} > 1)
\]

Precondition and postcondition assertions are presented in braces to distinguish them from parts of program statements. One possible precondition for this statement is \(\{x > 10\}\).

In axiomatic semantics, the meaning of a specific statement is defined by its precondition and its postcondition. In effect, the two assertions specify precisely the effect of executing the statement.
In the following subsections, we focus on correctness proofs of statements and programs, which is a common use of axiomatic semantics. The more general concept of axiomatic semantics is to state precisely the meaning of statements and programs in terms of logic expressions. Program verification is one application of axiomatic descriptions of languages.

### 3.5.3.2 Weakest Preconditions

The **weakest precondition** is the least restrictive precondition that will guarantee the validity of the associated postcondition. For example, in the statement and postcondition given in Section 3.5.3.1, \( \{x > 10\}, \{x > 50\}, \) and \( \{x > 1000\} \) are all valid preconditions. The weakest of all preconditions in this case is \( \{x > 0\} \).

If the weakest precondition can be computed from the most general postcondition for each of the statement types of a language, then the processes used to compute these preconditions provide a concise description of the semantics of that language. Furthermore, correctness proofs can be constructed for programs in that language. A program proof is begun by using the characteristics of the results of the program’s execution as the postcondition of the last statement of the program. This postcondition, along with the last statement, is used to compute the weakest precondition for the last statement. This precondition is then used as the postcondition for the second last statement. This process continues until the beginning of the program is reached. At that point, the precondition of the first statement states the conditions under which the program will compute the desired results. If these conditions are implied by the input specification of the program, the program has been verified to be correct.

An **inference rule** is a method of inferring the truth of one assertion on the basis of the values of other assertions. The general form of an inference rule is as follows:

\[
S_1, S_2, \ldots, S_n \\
\hline
S
\]

This rule states that if \( S_1, S_2, \ldots, \) and \( S_n \) are true, then the truth of \( S \) can be inferred. The top part of an inference rule is called its **antecedent**; the bottom part is called its **consequent**.

An **axiom** is a logical statement that is assumed to be true. Therefore, an axiom is an inference rule without an antecedent.

For some program statements, the computation of a weakest precondition from the statement and a postcondition is simple and can be specified by an axiom. In most cases, however, the weakest precondition can be specified only by an inference rule.

To use axiomatic semantics with a given programming language, whether for correctness proofs or for formal semantics specifications, either an axiom or an inference rule must exist for each kind of statement in the language.
In the following subsections, we present an axiom for assignment statements and inference rules for statement sequences, selection statements, and logical pretest loop statements. Note that we assume that neither arithmetic nor Boolean expressions have side effects.

### 3.5.3.3 Assignment Statements

The precondition and postcondition of an assignment statement together define its meaning. To define the meaning of an assignment statement there must be a way to compute its precondition from its postcondition.

Let $x = E$ be a general assignment statement and $Q$ be its postcondition. Then, its weakest precondition, $P$, is defined by the axiom

$$ P = Q_{x \rightarrow E} $$

which means that $P$ is computed as $Q$ with all instances of $x$ replaced by $E$. For example, if we have the assignment statement and postcondition

$$ a = b / 2 - 1 \{ a < 10 \} $$

the weakest precondition is computed by substituting $b / 2 - 1$ for $a$ in the postcondition $\{ a < 10 \}$, as follows:

$$ b / 2 - 1 < 10 \quad b < 22 $$

Thus, the weakest precondition for the given assignment statement and postcondition is $\{ b < 22 \}$. Remember that the assignment axiom is guaranteed to be correct only in the absence of side effects. An assignment statement has a side effect if it changes some variable other than its target.

The usual notation for specifying the axiomatic semantics of a given statement form is

$$ \{ P \} S \{ Q \} $$

where $P$ is the precondition, $Q$ is the postcondition, and $S$ is the statement form. In the case of the assignment statement, the notation is

$$ (Q_{x \rightarrow E}) \ x = E \{ Q \} $$

As another example of computing a precondition for an assignment statement, consider the following:

$$ x = 2 * y - 3 \{ x > 25 \} $$

The precondition is computed as follows:

$$ 2 * y - 3 > 25 \quad y > 14 $$
So \( \{ y > 14 \} \) is the weakest precondition for this assignment statement and postcondition.

Note that the appearance of the left side of the assignment statement in its right side does not affect the process of computing the weakest precondition. For example, for

\[
x = x + y - 3 \quad (x > 10)
\]

the weakest precondition is

\[
x + y - 3 > 10 \\
y > 13 - x
\]

Recall that axiomatic semantics was developed to prove the correctness of programs. In light of that, it is natural at this point to wonder how the axiom for assignment statements can be used to prove anything. Here is how: A given assignment statement with both a precondition and a postcondition can be considered a logical statement, or theorem. If the assignment axiom, when applied to the postcondition and the assignment statement, produces the given precondition, the theorem is proved. For example, consider the following logical statement:

\[
\{ x > 3 \} \quad x = x - 3 \quad (x > 0)
\]

Using the assignment axiom on the statement and its postcondition produces \( \{ x > 3 \} \), which is the given precondition. Therefore, we have proven the example logical statement.

Next, consider the following logical statement:

\[
\{ x > 5 \} \quad x = x - 3 \quad (x > 0)
\]

In this case, the given precondition, \( \{ x > 5 \} \), is not the same as the assertion produced by the axiom. However, it is obvious that \( \{ x > 5 \} \) implies \( \{ x > 3 \} \). To use this in a proof, an inference rule named the rule of consequence is needed. The form of the rule of consequence is

\[
\frac{P \Rightarrow P', Q \Rightarrow Q'}{P' \Rightarrow Q'}
\]

The \( \Rightarrow \) symbol means “implies,” and \( S \) can be any program statement. The rule can be stated as follows: If the logical statement \( \{ P \} S \{ Q \} \) is true, the assertion \( P' \) implies the assertion \( P \), and the assertion \( Q \) implies the assertion \( Q' \), then it can be inferred that \( \{ P' \} S \{ Q' \} \). In other words, the rule of consequence says that a postcondition can always be weakened and a precondition can always be strengthened. This is quite useful in program proofs. For example, it allows the completion of the proof of the last logical statement example above. If we let \( P \) be \( \{ x > 3 \} \), \( Q \) and \( Q' \) be \( \{ x > 0 \} \), and \( P' \) be \( \{ x > 5 \} \), we have
The first term of the antecedent \((x > 3) \quad x = x - 3 \quad (x > 0)\) was proven with the assignment axiom. The second and third terms are obvious. Therefore, by the rule of consequence, the consequent is true.

### 3.5.3.4 Sequences

The weakest precondition for a sequence of statements cannot be described by an axiom, because the precondition depends on the particular kinds of statements in the sequence. In this case, the precondition can only be described with an inference rule. Let \(S_1\) and \(S_2\) be adjacent program statements. If \(S_1\) and \(S_2\) have the following pre- and postconditions

\[
\{ P_1 \} \quad S_1 \quad \{ P_2 \}
\]

\[
\{ P_2 \} \quad S_2 \quad \{ P_3 \}
\]

the inference rule for such a two-statement sequence is

\[
\{ P_1 \} \quad S_1, S_2 \quad \{ P_3 \}
\]

So, for our example, \(\{ P_1 \} \quad S_1; S_2 \quad \{ P_2 \}\) describes the axiomatic semantics of the sequence \(S_1; S_2\). The inference rule states that to get the sequence precondition, the precondition of the second statement is computed. This new assertion is then used as the postcondition of the first statement, which can then be used to compute the precondition of the first statement, which is also the precondition of the whole sequence. If \(S_1\) and \(S_2\) are the assignment statements

\[
x_1 = E_1
\]

and

\[
x_2 = E_2
\]

then we have

\[
\{ P_3_{x_2 \to E_2} \} \quad x_2 = E_2 \quad \{ P_3 \}
\]

\[
\{ (P_3_{x_2 \to E_2})_{x_1 \to E_1} \} \quad x_1 = E_1 \quad \{ P_3_{x_2 \to E_2} \}
\]

Therefore, the weakest precondition for the sequence \(x_1 = E_1; x_2 = E_2\) with postcondition \(P_3\) is \(\{(P_3_{x_2 \to E_2})_{x_1 \to E_1}\}\).

For example, consider the following sequence and postcondition:

\[
y = 3 \ast x + 1;
\]

\[
x = y + 3;
\]

\[
\{ x < 10 \}
\]
The precondition for the second assignment statement is

\[ y < 7 \]

which is used as the postcondition for the first statement. The precondition for the first assignment statement can now be computed:

\[
3 \times x + 1 < 7 \\
x < 2
\]

So, \( x < 2 \) is the precondition of both the first statement and the two-statement sequence.

### 3.5.3.5 Selection

We next consider the inference rule for selection statements, the general form of which is

```
if B then S1 else S2
```

We consider only selections that include `else` clauses. The inference rule is

\[
\begin{align*}
\{ B \text{ and } P \} & \rightarrow \{Q\}, \\
\{ \text{not } B \text{ and } P \} & \rightarrow \{Q\}, \\
\{P\} & \rightarrow \{Q\}, \\
\end{align*}
\]

This rule specifies that selection statements must be proven both when the Boolean control expression is true and when it is false. The first logical statement above the line represents the `then` clause; the second represents the `else` clause. According to the inference rule, we need a precondition \( P \) that can be used in the precondition of both the `then` and `else` clauses.

Consider the following example of the computation of the precondition using the selection inference rule. The example selection statement is

```
if x > 0 then \\
y = y - 1 \\
else \\
y = y + 1
```

Suppose the postcondition, \( Q \), for this selection statement is \( \{y > 0\} \). We can use the axiom for assignment on the `then` clause

\[
y = y - 1 \ \{y > 0\}
\]

This produces \( \{y - 1 > 0\} \) or \( \{y > 1\} \). It can be used as the \( P \) part of the precondition for the `then` clause. Now we apply the same axiom to the `else` clause

\[
y = y + 1 \ \{y > 0\}
\]
which produces the precondition \(y + 1 > 0\) or \(y > -1\). Because \(y + 1 \Rightarrow y > -1\), the rule of consequence allows us to use \(y > 1\) for the precondition of the whole selection statement.

### 3.5.3.6 Logical Pretest Loops

Another essential construct of imperative programming languages is the logical pretest, or *while* loop. Computing the weakest precondition for a *while* loop is inherently more difficult than for a sequence, because the number of iterations cannot always be predetermined. In a case where the number of iterations is known, the loop can be unrolled and treated as a sequence.

The problem of computing the weakest precondition for loops is similar to the problem of proving a theorem about all positive integers. In the latter case, induction is normally used, and the same inductive method can be used for some loops. The principal step in induction is finding an inductive hypothesis. The corresponding step in the axiomatic semantics of a *while* loop is finding an assertion called a loop invariant, which is crucial to finding the weakest precondition.

The inference rule for computing the precondition for a *while* loop is as follows:

\[
\{I \text{ and } B\} \quad S \quad \{I\}
\]

\[
\{I\} \quad \text{while } B \quad \text{do} \quad S \quad \text{end} \quad \{I \text{ and } \neg B\}
\]

In this rule, \(I\) is the loop invariant. This seems simple, but it is not. The complexity lies in finding an appropriate loop invariant.

The axiomatic description of a *while* loop is written as

\[
\{P\} \quad \text{while } B \quad \text{do} \quad S \quad \text{end} \quad \{Q\}
\]

The loop invariant must satisfy a number of requirements to be useful. First, the weakest precondition for the *while* loop must guarantee the truth of the loop invariant. In turn, the loop invariant must guarantee the truth of the postcondition upon loop termination. These constraints move us from the inference rule to the axiomatic description. During execution of the loop, the truth of the loop invariant must be unaffected by the evaluation of the loop-controlling Boolean expression and the loop body statements. Hence, the name *invariant*.

Another complicating factor for *while* loops is the question of loop termination. A loop that does not terminate cannot be correct, and in fact computes nothing. If \(Q\) is the postcondition that holds immediately after loop exit, then a precondition \(P\) for the loop is one that guarantees \(Q\) at loop exit and also guarantees that the loop terminates.

The complete axiomatic description of a *while* construct requires all of the following to be true, in which \(I\) is the loop invariant:

\[
P \Rightarrow I
\]

\[
\{I \text{ and } B\} \quad S \quad \{I\}
\]

\[
(I \text{ and } \neg B)) \Rightarrow Q
\]

the loop terminates
If a loop computes a sequence of numeric values, it may be possible to find a loop invariant using an approach that is used for determining the inductive hypothesis when mathematical induction is used to prove a statement about a mathematical sequence. The relationship between the number of iterations and the precondition for the loop body is computed for a few cases, with the hope that a pattern emerges that will apply to the general case. It is helpful to treat the process of producing a weakest precondition as a function, \( \text{wp} \). In general:

\[
\text{wp}(\text{statement}, \text{postcondition}) = \text{precondition}
\]

A \( \text{wp} \) function is often called a **predicate transformer**, because it takes a predicate, or assertion, as a parameter and returns another predicate.

To find \( I \), the loop postcondition \( Q \) is used to compute preconditions for several different numbers of iterations of the loop body, starting with none. If the loop body contains a single assignment statement, the axiom for assignment statements can be used to compute these cases. Consider the example loop:

\[
\text{while } y <\!> x \text{ do } y = y + 1 \text{ end } (y = x)
\]

Remember that the equal sign is being used for two different purposes here. In assertions, it means mathematical equality; outside assertions, it means the assignment operator.

For zero iterations, the weakest precondition is, obviously, \( \{ y = x \} \).

For one iteration, it is

\[
\text{wp}(y = y + 1, \{ y = x \}) = \{ y + 1 = x \}, \text{ or } \{ y = x - 1 \}
\]

For two iterations, it is

\[
\text{wp}(y = y + 1, \{ y = x - 1 \}) = \{ y + 1 = x - 1 \}, \text{ or } \{ y = x - 2 \}
\]

For three iterations, it is

\[
\text{wp}(y = y + 1, \{ y = x - 2 \}) = \{ y + 1 = x - 2 \}, \text{ or } \{ y = x - 3 \}
\]

It is now obvious that \( \{ y < x \} \) will suffice for cases of one or more iterations. Combining this with \( \{ y = x \} \) for the zero iterations case, we get \( \{ y <= x \} \), which can be used for the loop invariant. A precondition for the \texttt{while} statement can be determined from the loop invariant. In fact, \( I \) can be used as the precondition, \( P \).

We must ensure that our choice satisfies the four criteria for \( I \) for our example loop. First, because \( P = I, P => I \). The second requirement is that the following must be true:

\[
\{ I \text{ and } B \} \text{ S } \{ I \}
\]

In our example, we have

\[
\{ y <= x \text{ and } y <\!> x \} \text{ y = y + 1 } \{ y <= x \}
\]
Applying the assignment axiom to

\[ y = y + 1 \{ y \leq x \} \]

we get \( \{ y + 1 \leq x \} \), which is equivalent to \( \{ y < x \} \), which is implied by \( \{ y \leq x \text{ and } y \neq x \} \). So, the earlier statement is proven.

Next, we must have

\( \{ I \text{ and } (\neg B) \} \Rightarrow Q \)

In our example, we have

\[
\begin{align*}
\{ (y \leq x) \text{ and } (\neg (y \neq x)) \} & \Rightarrow \{ y = x \} \\
\{ (y \leq x) \text{ and } (y = x) \} & \Rightarrow \{ y = x \} \\
\{ y = x \} & \Rightarrow \{ y = x \}
\end{align*}
\]

So, this is obviously true. Next, loop termination must be considered. In this example, the question is whether the loop

\( \{ y \leq x \} \text{ while } y \neq x \text{ do } y = y + 1 \text{ end } \{ y = x \} \)

terminates. Recalling that \( x \) and \( y \) are assumed to be integer variables, it is easy to see that this loop does terminate. The precondition guarantees that \( y \) initially is not larger than \( x \). The loop body increments \( y \) with each iteration, until \( y \) is equal to \( x \). No matter how much smaller \( y \) is than \( x \) initially, it will eventually become equal to \( x \). So the loop will terminate. Because our choice of \( I \) satisfies all four criteria, it is a satisfactory loop invariant and loop precondition.

The previous process used to compute the invariant for a loop does not always produce an assertion that is the weakest precondition (although it does in the example).

As another example of finding a loop invariant using the approach used in mathematical induction, consider the following loop statement:

\[
\text{while } s > 1 \text{ do } s = s / 2 \text{ end } \{ s = 1 \}
\]

As before, we use the assignment axiom to try to find a loop invariant and a precondition for the loop. For zero iterations, the weakest precondition is \( \{ s = 1 \} \). For one iteration, it is

\[
\text{wp}(s = s / 2, \{ s = 1 \}) = \{ s / 2 = 1 \}, \text{ or } \{ s = 2 \}
\]

For two iterations, it is

\[
\text{wp}(s = s / 2, \{ s = 2 \}) = \{ s / 2 = 2 \}, \text{ or } \{ s = 4 \}
\]

For three iterations, it is

\[
\text{wp}(s = s / 2, \{ s = 4 \}) = \{ s / 2 = 4 \}, \text{ or } \{ s = 8 \}
\]

From these cases, we can see clearly that the invariant is

\( \{ s \text{ is a nonnegative power of } 2 \} \)
Once again, the computed I can serve as P, and I passes the four requirements. Unlike our earlier example of finding a loop precondition, this one clearly is not a weakest precondition. Consider using the precondition \( \{ s > 1 \} \). The logical statement

\[
\{ s > 1 \} \textbf{ while } s > 1 \textbf{ do } s = s / 2 \textbf{ end } \{ s = 1 \}
\]

can easily be proven, and this precondition is significantly broader than the one computed earlier. The loop and precondition are satisfied for any positive value for \( s \), not just powers of 2, as the process indicates. Because of the rule of consequence, using a precondition that is stronger than the weakest precondition does not invalidate a proof.

Finding loop invariants is not always easy. It is helpful to understand the nature of these invariants. First, a loop invariant is a weakened version of the loop postcondition and also a precondition for the loop. So, I must be weak enough to be satisfied prior to the beginning of loop execution, but when combined with the loop exit condition, it must be strong enough to force the truth of the postcondition.

Because of the difficulty of proving loop termination, that requirement is often ignored. If loop termination can be shown, the axiomatic description of the loop is called \textbf{total correctness}. If the other conditions can be met but termination is not guaranteed, it is called \textbf{partial correctness}.

In more complex loops, finding a suitable loop invariant, even for partial correctness, requires a good deal of ingenuity. Because computing the precondition for a \textbf{while} loop depends on finding a loop invariant, proving the correctness of programs with \textbf{while} loops using axiomatic semantics can be difficult.

### 3.5.3.7 Program Proofs

This section provides validations for two simple programs. The first example of a correctness proof is for a very short program, consisting of a sequence of three assignment statements that interchange the values of two variables.

\[
\{ x = A \ \text{AND} \ y = B \}
\]
\[
t = x;
\]
\[
x = y;
\]
\[
y = t;
\]
\[
\{ x = B \ \text{AND} \ y = A \}
\]

Because the program consists entirely of assignment statements in a sequence, the assignment axiom and the inference rule for sequences can be used to prove its correctness. The first step is to use the assignment axiom on the last statement and the postcondition for the whole program. This yields the precondition

\[
\{ x = B \ \text{AND} \ t = A \}
\]
Next, we use this new precondition as a postcondition on the middle statement and compute its precondition, which is

\{ y = B \text{ AND } t = A \}\]

Next, we use this new assertion as the postcondition on the first statement and apply the assignment axiom, which yields

\{ y = B \text{ AND } x = A \}\]

which is the same as the precondition on the program, except for the order of operands on the \texttt{AND} operator. Because \texttt{AND} is a symmetric operator, our proof is complete.

The following example is a proof of correctness of a pseudocode program that computes the factorial function.

\{ n \geq 0 \}
count = n;
fact = 1;
while count \neq 0 do
    fact = fact \times count;
count = count - 1;
end
{fact = n!}

The method described earlier for finding the loop invariant does not work for the loop in this example. Some ingenuity is required here, which can be aided by a brief study of the code. The loop computes the factorial function in order of the last multiplication first; that is, \((n - 1) \times n\) is done first, assuming \(n\) is greater than 1. So, part of the invariant can be the following:

\text{fact} = (\text{count + 1}) \times (\text{count + 2}) \times \ldots \times (n - 1) \times n

But we also must ensure that \text{count} is always nonnegative, which we can do by adding that to the assertion above, to get the following:

\[ I = (\text{fact} = (\text{count + 1}) \times \ldots \times n) \text{ AND } (\text{count} \geq 0) \]

Next, we must confirm that this I meets the requirements for invariants. Once again we let I also be used for P, so P clearly implies I. The next question is

\{ I \text{ AND } B \} \Rightarrow \{ I \}

I and B is the following:

\((\text{fact} = (\text{count + 1}) \times \ldots \times n) \text{ AND } (\text{count} \geq 0)) \text{ AND } (\text{count} \neq 0)\]
This reduces to

\[(\text{fact} = (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count > 0)}\]

In our case, we must compute the precondition of the body of the loop, using the invariant for the postcondition. For

\[\{P\} \text{ count} = \text{count} - 1 \{I\}\]

we compute P to be

\[\{\text{(fact} = \text{count} \times (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 1)\}\]

Using this as the postcondition for the first assignment in the loop body,

\[\{P\} \text{ fact} = \text{fact} \times \text{count} \{\text{(fact} = \text{count} \times (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 1)\}\]

In this case, P is

\[\{\text{(fact} = (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 1)\}\]

It is clear that \(I\) and \(B\) implies this \(P\), so by the rule of consequence,

\[\{I \text{ AND } B\} S \{I\}\]

is true. Finally, the last test of \(I\) is

\(I \text{ AND (NOT }B) =\Rightarrow Q\)

For our example, this is

\[((\text{fact} = (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 0)) \Rightarrow \text{fact} = \text{n!}\]

This is clearly true, for when \(\text{count} = 0\), the first part is precisely the definition of factorial. So, our choice of \(I\) meets the requirements for a loop invariant. Now we can use our \(P\) (which is the same as \(I\)) from the \textbf{while} as the postcondition on the second assignment of the program

\[\{P\} \text{ fact} = 1 \{\text{(fact} = (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 0)\}\]

which yields for \(P\)

\[(1 = (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 0))\]

Using this as the postcondition for the first assignment in the code

\[\{P\} \text{ count} = \text{n} \{1 = (\text{count} + 1) \times \ldots \times \text{n}) \text{ AND (count }\geq 0)\}\]
produces for \( P \)

\[
((n + 1) \times \ldots \times n = 1) \land (n \geq 0)
\]

The left operand of the AND operator is true (because \( 1 = 1 \)) and the right operand is exactly the precondition of the whole code segment, \( \{n \geq 0\} \). Therefore, the program has been proven to be correct.

### 3.5.3.8 Evaluation

As stated previously, to define the semantics of a complete programming language using the axiomatic method, there must be an axiom or an inference rule for each statement type in the language. Defining axioms or inference rules for some of the statements of programming languages has proven to be a difficult task. An obvious solution to this problem is to design the language with the axiomatic method in mind, so that only statements for which axioms or inference rules can be written are included. Unfortunately, such a language would necessarily leave out some useful and powerful parts.

Axiomatic semantics is a powerful tool for research into program correctness proofs, and it provides an excellent framework in which to reason about programs, both during their construction and later. Its usefulness in describing the meaning of programming languages to language users and compiler writers is, however, highly limited.

### Summary

Backus-Naur Form and context-free grammars are equivalent metalanguages that are well suited for the task of describing the syntax of programming languages. Not only are they concise descriptive tools, but also the parse trees that can be associated with their generative actions give graphical evidence of the underlying syntactic structures. Furthermore, they are naturally related to recognition devices for the languages they generate, which leads to the relatively easy construction of syntax analyzers for compilers for these languages.

An attribute grammar is a descriptive formalism that can describe both the syntax and static semantics of a language. Attribute grammars are extensions to context-free grammars. An attribute grammar consists of a grammar, a set of attributes, a set of attribute computation functions, and a set of predicates that describe static semantics rules.

This chapter provides a brief introduction to three methods of semantic description: operational, denotational, and axiomatic. Operational semantics is a method of describing the meaning of language constructs in terms of their effects on an ideal machine. In denotational semantics, mathematical objects are used to represent the meanings of language constructs. Language entities are converted to these mathematical objects with recursive functions. Axiomatic semantics, which is based on formal logic, was devised as a tool for proving the correctness of programs.
Syntax description using context-free grammars and BNF are thoroughly discussed in Cleaveland and Uzgalis (1976).

Research in axiomatic semantics was begun by Floyd (1967) and further developed by Hoare (1969). The semantics of a large part of Pascal was described by Hoare and Wirth (1973) using this method. The parts they did not complete involved functional side effects and goto statements. These were found to be the most difficult to describe.

The technique of using preconditions and postconditions during the development of programs is described (and advocated) by Dijkstra (1976) and also discussed in detail in Gries (1981).

Good introductions to denotational semantics can be found in Gordon (1979) and Stoy (1977). Introductions to all of the semantics description methods discussed in this chapter can be found in Marcotty et al. (1976). Another good reference for much of the chapter material is Pagan (1981). The form of the denotational semantic functions in this chapter is similar to that found in Meyer (1990).

**Review Questions**

1. Define *lexeme* and *token*.
2. How are programming languages formally defined?
3. In which form is the programming language syntax commonly described?
4. What is a metalanguage?
5. What is a derivation in the context of grammar?
6. What is an ambiguous grammar?
7. What is a left-recursive grammar?
8. Explain the use of metasymbols in EBNFs.
9. What is the purpose of a predicate function?
10. When can the parse tree be called fully attributed?
11. How is the order of evaluation of attributes determined for the trees of a given attribute grammar?
12. What is the use of intrinsic attributes?
13. What is meant by decorating a parse tree?
14. Why can machine languages not be used to define statements in operational semantics?
15. Describe the two levels of uses of operational semantics.
16. In denotational semantics, what are the syntactic and semantic domains?
17. What is stored in the state of a program for denotational semantics?
18. What is an assertion in axiomatic semantics?
19. What two things must be defined for each language entity in order to construct a denotational description of the language?

20. Which part of an inference rule is the antecedent?

21. What is the weakest precondition?

22. What is an inference rule?

23. Give an example of a logical pretest loop.

24. On what branch of mathematics is denotational semantics based?

25. What is the problem with using a software pure interpreter for operational semantics?

26. Explain what the preconditions and postconditions of a given statement mean in axiomatic semantics.

27. What is the difference between total correctness and partial correctness with regard to loop termination?

28. When is a function called a predicate transformer?

29. In what fundamental way do operational semantics and denotational semantics differ?

---

1. The two mathematical models of language description are generation and recognition. Describe how each can define the syntax of a programming language.

2. Write EBNF descriptions for the following:
   a. A Java class definition header statement
   b. A Java method call statement
   c. A C \textbf{switch} statement
   d. A C \textbf{union} definition
   e. C \textbf{float} literals

3. Rewrite the BNF of Example 3.4 to give + precedence over * and force + to be right associative.

4. Rewrite the BNF of Example 3.4 to add the ++ and -- unary operators of Java.

5. Write a BNF description of the Boolean expressions of Java, including the three operators &&, ||, and ! and the relational expressions.

6. Using the grammar in Example 3.2, show a parse tree and a leftmost derivation for each of the following statements:
   a. \( A = A \times (B + (C \times A)) \)
   b. \( B = C \times (A \times C + B) \)
   c. \( A = A \times (B + (C)) \)
7. Using the grammar in Example 3.4, show a parse tree and a leftmost derivation for each of the following statements:
   a. \( A = (A + B) \times C \)
   b. \( A = B + C + A \)
   c. \( A = A \times (B + C) \)
   d. \( A = B \times (C \times (A + B)) \)

8. Prove that the following grammar is ambiguous:
   \[
   \begin{align*}
   &<S> \rightarrow <A> \\
   &<A> \rightarrow <A> + <A> \mid <id> \\
   &<id> \rightarrow a \mid b \mid c
   \end{align*}
   \]

9. Modify the grammar of Example 3.4 to add a unary minus operator that has higher precedence than either + or *.

10. Describe, in English, the language defined by the following grammar:
    \[
    \begin{align*}
    &<S> \rightarrow <A> <B> <C> \\
    &<A> \rightarrow a <A> \mid a \\
    &<B> \rightarrow b <B> \mid b \\
    &<C> \rightarrow \varepsilon
    \end{align*}
    \]

11. Consider the following grammar:
    \[
    \begin{align*}
    &<S> \rightarrow <A> a <B> b \\
    &<A> \rightarrow a <A> \mid b \\
    &<B> \rightarrow b
    \end{align*}
    \]
    Which of the following sentences are in the language generated by this grammar?
    a. babb
    b. bbbabb
    c. baaaaabc
    d. aaaaa

12. Consider the following grammar:
    \[
    \begin{align*}
    &<S> \rightarrow a <S> c <B> \mid <A> \\
    &<A> \rightarrow c <A> \mid c \\
    &<B> \rightarrow d
    \end{align*}
    \]
    Which of the following sentences are in the language generated by this grammar?
    a. abcd
    b. accbcd
    c. accbccc
    d. accd
    e. ccc
13. Write a grammar for the language consisting of strings that have \( n \) copies of the letter a followed by one more number of copies of the letter b, where \( n > 0 \). For example, the strings abb, aaaabbbbbb, and aaaaaaabbbbbbbbbbb are in the language but a, ab, ba, and aaabb are not.

14. Draw parse trees for the sentences abb and aabbb, as derived from the grammar of Problem 13.

15. Convert the BNF of Example 3.1 to EBNF.

16. Convert the BNF of Example 3.3 to EBNF.

17. Convert the following EBNF to BNF:
   \[
   S \rightarrow A \{ bA \} \\
   A \rightarrow a[b]A
   \]

18. What is the difference between an intrinsic attribute and a nonintrinsic synthesized attribute?

19. Write an attribute grammar whose BNF basis is that of Example 3.6 in Section 3.4.5 but whose language rules are as follows: Data types cannot be mixed in expressions, but assignment statements need not have the same types on both sides of the assignment operator.

20. Write an attribute grammar whose base BNF is that of Example 3.2 and whose type rules are the same as for the assignment statement example of Section 3.4.5.

21. Using the virtual machine instructions given in Section 3.5.1.1, give an operational semantic definition of the following:
   a. Java `do-while`
   b. Ada `for`
   c. C++ `if-then-else`
   d. C `for`
   e. C `switch`

22. Write a denotational semantics mapping function for the following statements:
   a. Ada `for`
   b. Java `do-while`
   c. Java Boolean expressions
   d. Java `for`
   e. C `switch`

23. Compute the weakest precondition for each of the following assignment statements and postconditions:
   a. \( a = 2 * (b - 1) - 1 \) \( \{ a > 0 \} \)
   b. \( b = (c + 10) / 3 \) \( \{ b > 6 \} \)
   c. \( a = a + 2 * b - 1 \) \( \{ a > 1 \} \)
   d. \( x = 2 * y + x - 1 \) \( \{ x > 11 \} \)
24. Compute the weakest precondition for each of the following sequences of assignment statements and their postconditions:

a. \( a = 2 \times b + 1; \)
   \( b = a - 3 \)
   \( \{ b < 0 \} \)

b. \( a = 3 \times (2 \times b + a); \)
   \( b = 2 \times a - 1 \)
   \( \{ b > 5 \} \)

25. Compute the weakest precondition for each of the following selection constructs and their postconditions:

a. \( \text{if } (a == b) \)
   \( b = 2 \times a + 1 \)
   \( \text{else} \)
   \( b = 2 \times a; \)
   \( \{ b > 1 \} \)

b. \( \text{if } (x < y) \)
   \( x = x + 1 \)
   \( \text{else} \)
   \( x = 3 \times x \)
   \( \{ x < 0 \} \)

c. \( \text{if } (x > y) \)
   \( y = 2 \times x + 1 \)
   \( \text{else} \)
   \( y = 3 \times x - 1; \)
   \( \{ y > 3 \} \)

26. Explain the four criteria for proving the correctness of a logical pretest loop construct of the form \textbf{while} B \textbf{do} S \textbf{end}.

27. Prove that \((n + 1) \times \ldots \times n = 1\).

28. Prove the following program is correct:

\[
\{ n > 0 \}
\text{count} = n;
\text{sum} = 0;
\textbf{while} \text{ count} <> 0 \textbf{ do }
\quad \text{sum} = \text{sum} + \text{count};
\quad \text{count} = \text{count} - 1;
\textbf{end}
\{ \text{sum} = 1 + 2 + \ldots + n \\}
4

Lexical and Syntax Analysis

4.1 Introduction
4.2 Lexical Analysis
4.3 The Parsing Problem
4.4 Recursive-Descent Parsing
4.5 Bottom-Up Parsing
This chapter begins with an introduction to lexical analysis, along with a simple example. Next, the general parsing problem is discussed, including the two primary approaches to parsing and the complexity of parsing. Then, we introduce the recursive-descent implementation technique for top-down parsers, including examples of parts of a recursive-descent parser and a trace of a parse using one. The last section discusses bottom-up parsing and the LR parsing algorithm. This section includes an example of a small LR parsing table and the parse of a string using the LR parsing process.

### 4.1 Introduction

A serious investigation of compiler design requires at least a semester of intensive study, including the design and implementation of a compiler for a small but realistic programming language. The first part of such a course is devoted to lexical and syntax analyses. The syntax analyzer is the heart of a compiler, because several other important components, including the semantic analyzer and the intermediate code generator, are driven by the actions of the syntax analyzer.

Some readers may wonder why a chapter on any part of a compiler would be included in a book on programming languages. There are at least two reasons to include a discussion of lexical and syntax analyses in this book: First, syntax analyzers are based directly on the grammars discussed in Chapter 3, so it is natural to discuss them as an application of grammars. Second, lexical and syntax analyzers are needed in numerous situations outside compiler design. Many applications, among them program listing formatters, programs that compute the complexity of programs, and programs that must analyze and react to the contents of a configuration file, all need to do lexical and syntax analyses. Therefore, lexical and syntax analyses are important topics for software developers, even if they never need to write a compiler. Furthermore, some computer science programs no longer require students to take a compiler design course, which leaves students with no instruction in lexical or syntax analysis. In those cases, this chapter can be covered in the programming language course. In degree programs that require a compiler design course, this chapter can be skipped.

Three different approaches to implementing programming languages are introduced in Chapter 1: compilation, pure interpretation, and hybrid implementation. The compilation approach uses a program called a compiler, which translates programs written in a high-level programming language into machine code. Compilation is typically used to implement programming languages that are used for large applications, often written in languages such as C++ and COBOL. Pure interpretation systems perform no translation; rather, programs are interpreted in their original form by a software interpreter. Pure interpretation is usually used for smaller systems in which execution efficiency is not critical, such as scripts embedded in HTML documents, written in languages such as JavaScript. Hybrid implementation systems translate programs written in high-level languages into intermediate forms, which are interpreted. These systems are now more widely used than ever, thanks in large part to the
4.2 Lexical Analysis

A lexical analyzer is essentially a pattern matcher. A pattern matcher attempts to find a substring of a given string of characters that matches a given character pattern. Pattern matching is a traditional part of computing. One of the earliest
uses of pattern matching was with text editors, such as the `ed` line editor, which
was introduced in an early version of UNIX. Since then, pattern matching
has found its way into some programming languages—for example, Perl and
JavaScript. It is also available through the standard class libraries of Java, C++,
and C#.

A lexical analyzer serves as the front end of a syntax analyzer. Technically,
lexical analysis is a part of syntax analysis. A lexical analyzer performs syntax
analysis at the lowest level of program structure. An input program appears to a
compiler as a single string of characters. The lexical analyzer collects characters
into logical groupings and assigns internal codes to the groupings according to
their structure. In Chapter 3, these logical groupings are named lexemes, and
the internal codes for categories of these groupings are named tokens. Lexemes
are recognized by matching the input character string against character string
patterns. Although tokens are usually represented as integer values, for the sake
of readability of lexical and syntax analyzers, they are often referenced through
named constants.

Consider the following example of an assignment statement:

```c
result = oldsum - value / 100;
```

Following are the tokens and lexemes of this statement:

<table>
<thead>
<tr>
<th>Token</th>
<th>Lexeme</th>
</tr>
</thead>
<tbody>
<tr>
<td>IDENT</td>
<td>result</td>
</tr>
<tr>
<td>ASSIGN_OP</td>
<td>=</td>
</tr>
<tr>
<td>IDENT</td>
<td>oldsum</td>
</tr>
<tr>
<td>SUB_OP</td>
<td>-</td>
</tr>
<tr>
<td>IDENT</td>
<td>value</td>
</tr>
<tr>
<td>DIV_OP</td>
<td>/</td>
</tr>
<tr>
<td>INT_LIT</td>
<td>100</td>
</tr>
<tr>
<td>SEMICOLON</td>
<td>;</td>
</tr>
</tbody>
</table>

Lexical analyzers extract lexemes from a given input string and produce the
corresponding tokens. In the early days of compilers, lexical analyzers often
processed an entire source program file and produced a file of tokens and lexemes. Now, however, most lexical analyzers are subprograms that locate the
next lexeme in the input, determine its associated token code, and return them
to the caller, which is the syntax analyzer. So, each call to the lexical analyzer
returns a single lexeme and its token. The only view of the input program seen
by the syntax analyzer is the output of the lexical analyzer, one token at a time.

The lexical-analysis process includes skipping comments and white space
outside lexemes, as they are not relevant to the meaning of the program. Also,
the lexical analyzer inserts lexemes for user-defined names into the symbol
table, which is used by later phases of the compiler. Finally, lexical analyzers
detect syntactic errors in tokens, such as ill-formed floating-point literals, and
report such errors to the user.
There are three approaches to building a lexical analyzer:

1. Write a formal description of the token patterns of the language using a descriptive language related to regular expressions.¹ These descriptions are used as input to a software tool that automatically generates a lexical analyzer. There are many such tools available for this. The oldest of these, named lex, is commonly included as part of UNIX systems.

2. Design a state transition diagram that describes the token patterns of the language and write a program that implements the diagram.

3. Design a state transition diagram that describes the token patterns of the language and hand-construct a table-driven implementation of the state diagram.

A state transition diagram, or just state diagram, is a directed graph. The nodes of a state diagram are labeled with state names. The arcs are labeled with the input characters that cause the transitions among the states. An arc may also include actions the lexical analyzer must perform when the transition is taken.

State diagrams of the form used for lexical analyzers are representations of a class of mathematical machines called finite automata. Finite automata can be designed to recognize members of a class of languages called regular languages. Regular grammars are generative devices for regular languages. The tokens of a programming language are a regular language, and a lexical analyzer is a finite automaton.

We now illustrate lexical-analyzer construction with a state diagram and the code that implements it. The state diagram could simply include states and transitions for each and every token pattern. However, that approach results in a very large and complex diagram, because every node in the state diagram would need a transition for every character in the character set of the language being analyzed. We therefore consider ways to simplify it.

Suppose we need a lexical analyzer that recognizes only arithmetic expressions, including variable names and integer literals as operands. Assume that the variable names consist of strings of uppercase letters, lowercase letters, and digits but must begin with a letter. Names have no length limitation. The first thing to observe is that there are 52 different characters (any uppercase or lowercase letter) that can begin a name, which would require 52 transitions from the transition diagram’s initial state. However, a lexical analyzer is interested only in determining that it is a name and is not concerned with which specific name it happens to be. Therefore, we define a character class named LETTER for all 52 letters and use a single transition on the first letter of any name.

Another opportunity for simplifying the transition diagram is with the integer literal tokens. There are 10 different characters that could begin an integer literal lexeme. This would require 10 transitions from the start state of the state diagram. Because specific digits are not a concern of the lexical

¹. These regular expressions are the basis for the pattern-matching facilities now part of many programming languages, either directly or through a class library.
analyzer, we can build a much more compact state diagram if we define a character class named DIGIT for digits and use a single transition on any character in this character class to a state that collects integer literals.

Because our names can include digits, the transition from the node following the first character of a name can use a single transition on LETTER or DIGIT to continue collecting the characters of a name.

Next, we define some utility subprograms for the common tasks inside the lexical analyzer. First, we need a subprogram, which we can name getChar, that has several duties. When called, getChar gets the next character of input from the input program and puts it in the global variable nextChar. getChar also must determine the character class of the input character and put it in the global variable charClass. The lexeme being built by the lexical analyzer, which could be implemented as a character string or an array, will be named lexeme.

We implement the process of putting the character in nextChar into the string array lexeme in a subprogram named addChar. This subprogram must be explicitly called because programs include some characters that need not be put in lexeme, for example the white-space characters between lexemes. In a more realistic lexical analyzer, comments also would not be placed in lexeme.

When the lexical analyzer is called, it is convenient if the next character of input is the first character of the next lexeme. Because of this, a function named getNonBlank is used to skip white space every time the analyzer is called.

Finally, a subprogram named lookup is needed to compute the token code for the single-character tokens. In our example, these are parentheses and the arithmetic operators. Token codes are numbers arbitrarily assigned to tokens by the compiler writer.

The state diagram in Figure 4.1 describes the patterns for our tokens. It includes the actions required on each transition of the state diagram.

The following is a C implementation of a lexical analyzer specified in the state diagram of Figure 4.1, including a main driver function for testing purposes:

```c
/* front.c - a lexical analyzer system for simple arithmetic expressions */

#include <stdio.h>
#include <ctype.h>

/* Global declarations */
/* Variables */
int charClass;
char lexeme [100];
char nextChar;
int lexLen;
int token;
int nextToken;
FILE *in_fp, *fopen();
```
/* Function declarations */
void addChar();
void getChar();
void getNonBlank();
int lex();

/* Character classes */
#define LETTER 0
#define DIGIT 1
#define UNKNOWN 99

/* Token codes */
#define INT_LIT 10
#define IDENT 11
#define ASSIGN_OP 20
#define ADD_OP 21
#define SUB_OP 22
#define MULT_OP 23
#define DIV_OP 24
#define LEFT_PAREN 25
#define RIGHT_PAREN 26
Chapter 4  Lexical and Syntax Analysis

/****************************************************************************
 /* main driver */
 main() {

 /* Open the input data file and process its contents */
 if ((in_fp = fopen("front.in", "r")) == NULL)
   printf("ERROR - cannot open front.in \n");
 else {
   getChar();
   do {
     lex();
   } while (nextToken != EOF);
 }
}

/****************************************************************************
 /* lookup - a function to lookup operators and parentheses and return the token */
 int lookup(char ch) {
 switch (ch) {
 case '(':
   addChar();
   nextToken = LEFT_PAREN;
   break;

 case ')':
   addChar();
   nextToken = RIGHT_PAREN;
   break;

 case '+':
   addChar();
   nextToken = ADD_OP;
   break;

 case '-':
   addChar();
   nextToken = SUB_OP;
   break;

 case '*':
   addChar();
   nextToken = MULT_OP;
   break;

 case '/':
   addChar();
   nextToken = DIV_OP;
   break;

 /* Other cases */
 }
 return nextToken;
}
4.2 Lexical Analysis

```c
default:
    addChar();
    nextToken = EOF;
    break;
}
return nextToken;
}

/****************************************************************************
/* addChar - a function to add nextChar to lexeme */
void addChar() {
    if (lexLen <= 98) {
        lexeme[lexLen++] = nextChar;
        lexeme[lexLen] = 0;
    } else
        printf("Error - lexeme is too long \n");
}

/****************************************************************************
/* getChar - a function to get the next character of input and determine its character class */
void getChar() {
    if ((nextChar = getc(in_fp)) != EOF) {
        if (isalpha(nextChar))
            charClass = LETTER;
        else if (isdigit(nextChar))
            charClass = DIGIT;
        else charClass = UNKNOWN;
    } else
        charClass = EOF;
}

/****************************************************************************
/* getNonBlank - a function to call getChar until it returns a non-whitespace character */
void getNonBlank() {
    while (isspace(nextChar))
        getChar();
}

/****************************************************************************
/* lex - a simple lexical analyzer for arithmetic expressions */
int lex() {
    lexLen = 0;
    getNonBlank();
    switch (charClass) {
```
Chapter 4  Lexical and Syntax Analysis

/* Parse identifiers */
case LETTER:
    addChar();
    getChar();
    while (charClass == LETTER || charClass == DIGIT) {
        addChar();
        getChar();
    }
    nextToken = IDENT;
    break;

/* Parse integer literals */
case DIGIT:
    addChar();
    getChar();
    while (charClass == DIGIT) {
        addChar();
        getChar();
    }
    nextToken = INT_LIT;
    break;

/* Parentheses and operators */
case UNKNOWN:
    lookup(nextChar);
    getChar();
    break;

/* EOF */
case EOF:
    nextToken = EOF;
    lexeme[0] = 'E';
    lexeme[1] = 'O';
    lexeme[2] = 'F';
    lexeme[3] = 0;
    break;
} /* End of switch */
printf("Next token is: %d, Next lexeme is %s\n", 
    nextToken, lexeme);
return nextToken;
} /* End of function lex */

This code illustrates the relative simplicity of lexical analyzers. Of course, we have left out input buffering, as well as some other important details. Furthermore, we have dealt with a very small and simple input language.

Consider the following expression:

(sum + 47) / total
Following is the output of the lexical analyzer of `front.c` when used on this expression:

```plaintext
Next token is: 25 Next lexeme is (
Next token is: 11 Next lexeme is sum
Next token is: 21 Next lexeme is +
Next token is: 10 Next lexeme is 47
Next token is: 26 Next lexeme is )
Next token is: 24 Next lexeme is /
Next token is: 11 Next lexeme is total
Next token is: -1 Next lexeme is EOF
```

Names and reserved words in programs have similar patterns. Although it is possible to build a state diagram to recognize every specific reserved word of a programming language, that would result in a prohibitively large state diagram. It is much simpler and faster to have the lexical analyzer recognize names and reserved words with the same pattern and use a lookup in a table of reserved words to determine which names are reserved words. Using this approach considers reserved words to be exceptions in the names token category.

A lexical analyzer often is responsible for the initial construction of the symbol table, which acts as a database of names for the compiler. The entries in the symbol table store information about user-defined names, as well as the attributes of the names. For example, if the name is that of a variable, the variable's type is one of its attributes that will be stored in the symbol table. Names are usually placed in the symbol table by the lexical analyzer. The attributes of a name are usually put in the symbol table by some part of the compiler that is subsequent to the actions of the lexical analyzer.

### 4.3 The Parsing Problem

The part of the process of analyzing syntax that is referred to as syntax analysis is often called parsing. We will use these two interchangeably.

This section discusses the general parsing problem and introduces the two main categories of parsing algorithms, top-down and bottom-up, as well as the complexity of the parsing process.

#### 4.3.1 Introduction to Parsing

Parsers for programming languages construct parse trees for given programs. In some cases, the parse tree is only implicitly constructed, meaning that perhaps only a traversal of the tree is generated. But in all cases, the information required to build the parse tree is created during the parse. Both parse trees and derivations include all of the syntactic information needed by a language processor.
There are two distinct goals of syntax analysis: First, the syntax analyzer must check the input program to determine whether it is syntactically correct. When an error is found, the analyzer must produce a diagnostic message and recover. In this case, recovery means it must get back to a normal state and continue its analysis of the input program. This step is required so that the compiler finds as many errors as possible during a single analysis of the input program. If it is not done well, error recovery may create more errors, or at least more error messages. The second goal of syntax analysis is to produce a complete parse tree, or at least trace the structure of the complete parse tree, for syntactically correct input. The parse tree (or its trace) is used as the basis for translation.

Parsers are categorized according to the direction in which they build parse trees. The two broad classes of parsers are top-down, in which the tree is built from the root downward to the leaves, and bottom-up, in which the parse tree is built from the leaves upward to the root.

In this chapter, we use a small set of notational conventions for grammar symbols and strings to make the discussion less cluttered. For formal languages, they are as follows:

1. Terminal symbols—lowercase letters at the beginning of the alphabet (a, b, . . .)
2. Nonterminal symbols—uppercase letters at the beginning of the alphabet (A, B, . . .)
3. Terminals or nonterminals—uppercase letters at the end of the alphabet (W, X, Y, Z)
4. Strings of terminals—lowercase letters at the end of the alphabet (w, x, y, z)
5. Mixed strings (terminals and/or nonterminals)—lowercase Greek letters (α, β, δ, γ)

For programming languages, terminal symbols are the small-scale syntactic constructs of the language, what we have referred to as lexemes. The nonterminal symbols of programming languages are usually connotative names or abbreviations, surrounded by angle brackets—for example, <while_statement>, <expr>, and <function_def>. The sentences of a language (programs, in the case of a programming language) are strings of terminals. Mixed strings describe right-hand sides (RHSs) of grammar rules and are used in parsing algorithms.

### 4.3.2 Top-Down Parsers

A top-down parser traces or builds a parse tree in preorder. A preorder traversal of a parse tree begins with the root. Each node is visited before its branches are followed. Branches from a particular node are followed in left-to-right order. This corresponds to a leftmost derivation.
In terms of the derivation, a top-down parser can be described as follows: Given a sentential form that is part of a leftmost derivation, the parser’s task is to find the next sentential form in that leftmost derivation. The general form of a left sentential form is $xA\alpha$, whereby our notational conventions $x$ is a string of terminal symbols, $A$ is a nonterminal, and $\alpha$ is a mixed string. Because $x$ contains only terminals, $A$ is the leftmost nonterminal in the sentential form, so it is the one that must be expanded to get the next sentential form in a leftmost derivation. Determining the next sentential form is a matter of choosing the correct grammar rule that has $A$ as its LHS. For example, if the current sentential form is $xA\alpha$ and the $A$-rules are $A \rightarrow bB$, $A \rightarrow cBb$, and $A \rightarrow a$, a top-down parser must choose among these three rules to get the next sentential form, which could be $xbB\alpha$, $xcBb\alpha$, or $xa\alpha$. This is the parsing decision problem for top-down parsers.

Different top-down parsing algorithms use different information to make parsing decisions. The most common top-down parsers choose the correct RHS for the leftmost nonterminal in the current sentential form by comparing the next token of input with the first symbols that can be generated by the RHSs of those rules. Whichever RHS has that token at the left end of the string it generates is the correct one. So, in the sentential form $xA\alpha$, the parser would use whatever token would be the first generated by $A$ to determine which $A$-rule should be used to get the next sentential form. In the example above, the three RHSs of the $A$-rules all begin with different terminal symbols. The parser can easily choose the correct RHS based on the next token of input, which must be $a$, $b$, or $c$ in this example. In general, choosing the correct RHS is not so straightforward, because some of the RHSs of the leftmost nonterminal in the current sentential form may begin with a nonterminal.

The most common top-down parsing algorithms are closely related. A **recursive-descent parser** is a coded version of a syntax analyzer based directly on the BNF description of the syntax of language. The most common alternative to recursive descent is to use a parsing table, rather than code, to implement the BNF rules. Both of these, which are called LL algorithms, are equally powerful, meaning they work on the same subset of all context-free grammars. The first L in LL specifies a left-to-right scan of the input; the second L specifies that a leftmost derivation is generated. Section 4.4 introduces the recursive-descent approach to implementing an LL parser.

### 4.3.3 Bottom-Up Parsers

A bottom-up parser constructs a parse tree by beginning at the leaves and progressing toward the root. This parse order corresponds to the reverse of a rightmost derivation. That is, the sentential forms of the derivation are produced in order of last to first. In terms of the derivation, a bottom-up parser can be described as follows: Given a right sentential form $\alpha$, the parser must determine what substring of $\alpha$ is the RHS of the rule in the grammar that must be reduced to its LHS to produce the previous sentential form in the rightmost
derivation. For example, the first step for a bottom-up parser is to determine which substring of the initial given sentence is the RHS to be reduced to its corresponding LHS to get the second last sentential form in the derivation. The process of finding the correct RHS to reduce is complicated by the fact that a given right sentential form may include more than one RHS from the grammar of the language being parsed. The correct RHS is called the handle. A right sentential form is a sentential form that appears in a rightmost derivation.

Consider the following grammar and derivation:

\[
\begin{align*}
S & \rightarrow aAc \\
A & \rightarrow aA \mid b \\
S & \rightarrow aAc \Rightarrow aaAc \Rightarrow aabc
\end{align*}
\]

A bottom-up parser of this sentence, aabc, starts with the sentence and must find the handle in it. In this example, this is an easy task, for the string contains only one RHS, b. When the parser replaces b with its LHS, A, it gets the second to last sentential form in the derivation, aaAc. In the general case, as stated previously, finding the handle is much more difficult, because a sentential form may include several different RHSs.

A bottom-up parser finds the handle of a given right sentential form by examining the symbols on one or both sides of a possible handle. Symbols to the right of the possible handle are usually tokens in the input that have not yet been analyzed.

The most common bottom-up parsing algorithms are in the LR family, where the L specifies a left-to-right scan of the input and the R specifies that a rightmost derivation is generated.

### 4.3.4 The Complexity of Parsing

Parsing algorithms that work for any unambiguous grammar are complicated and inefficient. In fact, the complexity of such algorithms is \(O(n^3)\), which means the amount of time they take is on the order of the cube of the length of the string to be parsed. This relatively large amount of time is required because these algorithms frequently must back up and reparse part of the sentence being analyzed. Reparsing is required when the parser has made a mistake in the parsing process. Backing up the parser also requires that part of the parse tree being constructed (or its trace) must be dismantled and rebuilt. \(O(n^3)\) algorithms are normally not useful for practical processes, such as syntax analysis for a compiler, because they are far too slow. In situations such as this, computer scientists often search for algorithms that are faster, though less general. Generality is traded for efficiency. In terms of parsing, faster algorithms have been found that work for only a subset of the set of all possible grammars. These algorithms are acceptable as long as the subset includes grammars that describe programming languages. (Actually, as discussed in Chapter 3, the whole class
of context-free grammars is not adequate to describe all of the syntax of most programming languages.)

All algorithms used for the syntax analyzers of commercial compilers have complexity $O(n)$, which means the time they take is linearly related to the length of the string to be parsed. This is vastly more efficient than $O(n^3)$ algorithms.

### 4.4 Recursive-Descent Parsing

This section introduces the recursive-descent top-down parser implementation process.

#### 4.4.1 The Recursive-Descent Parsing Process

A recursive-descent parser is so named because it consists of a collection of subprograms, many of which are recursive, and it produces a parse tree in top-down order. This recursion is a reflection of the nature of programming languages, which include several different kinds of nested structures. For example, statements are often nested in other statements. Also, parentheses in expressions must be properly nested. The syntax of these structures is naturally described with recursive grammar rules.

EBNF is ideally suited for recursive-descent parsers. Recall from Chapter 3 that the primary EBNF extensions are braces, which specify that what they enclose can appear zero or more times, and brackets, which specify that what they enclose can appear once or not at all. Note that in both cases, the enclosed symbols are optional. Consider the following examples:

```plaintext
<if_statement> → if <logic_expr> <statement> [else <statement>]
<ident_list> → ident {, ident}
```

In the first rule, the `else` clause of an `if` statement is optional. In the second, an `<ident_list>` is an identifier, followed by zero or more repetitions of a comma and an identifier.

A recursive-descent parser has a subprogram for each nonterminal in its associated grammar. The responsibility of the subprogram associated with a particular nonterminal is as follows: When given an input string, it traces out the parse tree that can be rooted at that nonterminal and whose leaves match the input string. In effect, a recursive-descent parsing subprogram is a parser for the language (set of strings) that is generated by its associated nonterminal.

Consider the following EBNF description of simple arithmetic expressions:

```plaintext
<expr> → <term> {((+ | -) <term>)}
<term> → <factor> {((* | /) <factor>)}
<factor> → id | int_constant | ( <expr> )
```
Recall from Chapter 3 that an EBNF grammar for arithmetic expressions, such as this one, does not force any associativity rule. Therefore, when using such a grammar as the basis for a compiler, one must take care to ensure that the code generation process, which is normally driven by syntax analysis, produces code that adheres to the associativity rules of the language. This can be done easily when recursive-descent parsing is used.

In the following recursive-descent function, `expr`, the lexical analyzer is the function that is implemented in Section 4.2. It gets the next lexeme and puts its token code in the global variable `nextToken`. The token codes are defined as named constants, as in Section 4.2.

A recursive-descent subprogram for a rule with a single RHS is relatively simple. For each terminal symbol in the RHS, that terminal symbol is compared with `nextToken`. If they do not match, it is a syntax error. If they match, the lexical analyzer is called to get the next input token. For each nonterminal, the parsing subprogram for that nonterminal is called.

The recursive-descent subprogram for the first rule in the previous example grammar, written in C, is

```c
/* expr
   Parses strings in the language generated by the rule:
   <expr> -> <term> {(+ | -) <term>}
*/
void expr() {
    printf("Enter <expr>n");

    /* Parse the first term */
    term();

    /* As long as the next token is + or -, get
        the next token and parse the next term */
    while (nextToken == ADD_OP || nextToken == SUB_OP) {
        lex();
        term();
    }
    printf("Exit <expr>n");
}  /* End of function expr */
```

Notice that the `expr` function includes tracing output statements, which are included to produce the example output shown later in this section.

Recursive-descent parsing subprograms are written with the convention that each one leaves the next token of input in `nextToken`. So, whenever a parsing function begins, it assumes that `nextToken` has the code for the leftmost token of the input that has not yet been used in the parsing process.

The part of the language that the `expr` function parses consists of one or more terms, separated by either plus or minus operators. This is the language generated by the nonterminal `<expr>`. Therefore, first it calls the function that parses terms (`term`). Then it continues to call that function as long as
it finds ADD_OP or SUB_OP tokens (which it passes over by calling lex). This recursive-descent function is simpler than most, because its associated rule has only one RHS. Furthermore, it does not include any code for syntax error detection or recovery, because there are no detectable errors associated with the grammar rule.

A recursive-descent parsing subprogram for a nonterminal whose rule has more than one RHS begins with code to determine which RHS is to be parsed. Each RHS is examined (at compiler construction time) to determine the set of terminal symbols that can appear at the beginning of sentences it can generate. By matching these sets against the next token of input, the parser can choose the correct RHS.

The parsing subprogram for <term> is similar to that for <expr>:

```c
/* term
   Parses strings in the language generated by the rule:
   <term> -> <factor> {(* | /) <factor>}
*/
void term() {
    printf("Enter <term>\n");

    /* Parse the first factor */
    factor();

    /* As long as the next token is * or /, get the next token and parse the next factor */
    while (nextToken == MULT_OP || nextToken == DIV_OP) {
        lex();
        factor();
    }
    printf("Exit <term>\n"); /* End of function term */
}
```

The function for the <factor> nonterminal of our arithmetic expression grammar must choose between its two RHSs. It also includes error detection. In the function for <factor>, the reaction to detecting a syntax error is simply to call the error function. In a real parser, a diagnostic message must be produced when an error is detected. Furthermore, parsers must recover from the error so that the parsing process can continue.

```c
/* factor
   Parses strings in the language generated by the rule:
   <factor> -> id | int_constant | ( <expr >
*/
void factor() {
    printf("Enter <factor>\n");

    /* Determine which RHS */
    if (nextToken == IDENT || nextToken == INT_LIT)
```
/* Get the next token */
    lex();

/* If the RHS is ( <expr>), call lex to pass over the
left parenthesis, call expr, and check for the right
parenthesis */
else {
    if (nextToken == LEFT_PAREN) {
        lex();
        expr();
        if (nextToken == RIGHT_PAREN)
            lex();
        else
            error();
    } /* End of if (nextToken == ... */

/* It was not an id, an integer literal, or a left
parenthesis */
else
    error();
} /* End of else */

printf("Exit <factor>\n");
} /* End of function factor */

Following is the trace of the parse of the example expression (sum + 47) /
total, using the parsing functions expr, term, and factor, and the function
lex from Section 4.2. Note that the parse begins by calling lex and the start
symbol routine, in this case, expr.

Next token is: 25 Next lexeme is ( Enter <expr> Enter <term> Enter <factor>
Next token is: 11 Next lexeme is sum Enter <expr> Enter <term> Enter <factor>
Next token is: 21 Next lexeme is + Exit <factor> Exit <term> Exit <expr>
Next token is: 10 Next lexeme is 47 Enter <term> Enter <factor> Next token is: 26 Next lexeme is ) Exit <factor> Exit <term> Exit <expr>
Next token is: 24 Next lexeme is /
Exit <factor>
Next token is: 11 Next lexeme is total
Enter <factor>
Next token is: -1 Next lexeme is EOF
Exit <factor>
Exit <term>
Exit <expr>

The parse tree traced by the parser for the preceding expression is shown in Figure 4.2.

Figure 4.2
Parse tree for

\[
\frac{\text{sum} + 47}{\text{total}}
\]

One more example grammar rule and parsing function should help solidify the reader’s understanding of recursive-descent parsing. Following is a grammatical description of the Java if statement:

\[
<\text{ifstmt}> \rightarrow \text{if} \ (<\text{boolexpr}>\) <\text{statement}> [\text{else} <\text{statement}>]
\]

The recursive-descent subprogram for this rule follows:

```c
/* Function ifstmt 
Parses strings in the language generated by the rule:
<ifstmt> -> if (<boolexpr>) <statement> 
[else <statement>] */

void ifstmt() {
/* Be sure the first token is 'if' */
    if (nextToken != IF_CODE)
        error();
    else {
```
/* Call lex to get to the next token */
lex();
/* Check for the left parenthesis */
if (nextToken != LEFT_PAREN)
    error();
else {
/* Parse the Boolean expression */
    boolexpr();
/* Check for the right parenthesis */
    if (nextToken != RIGHT_PAREN)
        error();
    else {
/* Parse the then clause */
        statement();
/* If an else is next, parse the else clause */
    if (nextToken == ELSE_CODE) {
/* Call lex to get over the else */
        lex();
        statement();
    } /* end of if (nextToken == ELSE_CODE ...) */
    } /* end of else of if (nextToken != RIGHT ... */
    } /* end of else of if (nextToken != IF_CODE ... */
} /* end of ifstmt */

Notice that this function uses parser functions for statements and Boolean expressions that are not given in this section.

The objective of these examples is to convince you that a recursive-descent parser can be easily written if an appropriate grammar is available for the language. The characteristics of a grammar that allows a recursive-descent parser to be built are discussed in the following subsection.

### 4.4.2 The LL Grammar Class

Before choosing to use recursive descent as a parsing strategy for a compiler or other program analysis tool, one must consider the limitations of the approach, in terms of grammar restrictions. This section discusses these restrictions and their possible solutions.

One simple grammar characteristic that causes a catastrophic problem for LL parsers is left recursion. For example, consider the following rule:

\[ A \rightarrow A + B \]

A recursive-descent parser subprogram for A immediately calls itself to parse the first symbol in its RHS. That activation of the A parser subprogram then immediately calls itself again, and again, and so forth. It is easy to see that this leads nowhere (except to stack overflow).

The left recursion in the rule \( A \rightarrow A + B \) is called **direct left recursion**, because it occurs in one rule. Direct left recursion can be eliminated from a grammar by the following process:
For each nonterminal, A,

1. Group the A-rules as

   \[ A \rightarrow A_{1}, \ldots, A_{m} \mid \beta_{1} \mid \beta_{2} \mid \ldots \mid \beta_{n} \]

   where none of the \( \beta \)'s begins with A

2. Replace the original A-rules with

   \[ A \rightarrow \beta_{1}A' \mid \beta_{2}A' \mid \ldots \mid \beta_{n}A' \]

   \[ A' \rightarrow \alpha_{1}A' \mid \alpha_{2}A' \mid \alpha_{m}A' \mid \varepsilon \]

Note that \( \varepsilon \) specifies the empty string. A rule that has \( \varepsilon \) as its RHS is called an erasure rule, because its use in a derivation effectively erases its LHS from the sentential form.

Consider the following example grammar and the application of the above process:

\[
\begin{align*}
E & \rightarrow E + T \mid T \\
T & \rightarrow T * F \mid F \\
F & \rightarrow (E) \mid id
\end{align*}
\]

For the E-rules, we have \( \alpha_{1} = + \ T \) and \( \beta = T \), so we replace the E-rules with

\[
\begin{align*}
E & \rightarrow T E' \\
E' & \rightarrow + T E' \mid \varepsilon
\end{align*}
\]

For the T-rules, we have \( \alpha_{1} = *F \) and \( \beta = F \), so we replace the T-rules with

\[
\begin{align*}
T & \rightarrow F T' \\
T' & \rightarrow * F T' \mid \varepsilon
\end{align*}
\]

Because there is no left recursion in the F-rules, they remain the same, so the complete replacement grammar is

\[
\begin{align*}
E & \rightarrow T E' \\
E' & \rightarrow + T E' \mid \varepsilon \\
T & \rightarrow F T' \\
T' & \rightarrow * F T' \mid \varepsilon \\
F & \rightarrow (E) \mid id
\end{align*}
\]

This grammar generates the same language as the original grammar but is not left recursive.

As was the case with the expression grammar written using EBNF in Section 4.4.1, this grammar does not specify left associativity of operators. However, it is relatively easy to design the code generation based on this grammar so that the addition and multiplication operators will have left associativity.

Indirect left recursion poses the same problem as direct left recursion. For example, suppose we have

\[
\begin{align*}
A & \rightarrow B \ a \ A \\
B & \rightarrow A \ b
\end{align*}
\]
A recursive-descent parser for these rules would have the A subprogram immediately call the subprogram for B, which immediately calls the A subprogram. So, the problem is the same as for direct left recursion. The problem of left recursion is not confined to the recursive-descent approach to building top-down parsers. It is a problem for all top-down parsing algorithms. Fortunately, left recursion is not a problem for bottom-up parsing algorithms.

There is an algorithm to modify a given grammar to remove indirect left recursion (Aho et al., 2006), but it is not covered here. When writing a grammar for a programming language, one can usually avoid including left recursion, both direct and indirect.

Left recursion is not the only grammar trait that disallows top-down parsing. Another is whether the parser can always choose the correct RHS on the basis of the next token of input, using only the first token generated by the leftmost nonterminal in the current sentential form. There is a relatively simple test of a non-left recursive grammar that indicates whether this can be done, called the pairwise disjointness test. This test requires the ability to compute a set based on the RHSs of a given nonterminal symbol in a grammar. These sets, which are called FIRST, are defined as

\[
\text{FIRST}(\alpha) = \{a \mid \alpha \Rightarrow^* a\beta\} \quad \text{(If } \alpha \Rightarrow^* e, e \text{ is in FIRST}(\alpha))
\]

in which \(\Rightarrow^*\) means 0 or more derivation steps.

An algorithm to compute FIRST for any mixed string \(\alpha\) can be found in Aho et al. (2006). For our purposes, FIRST can usually be computed by inspection of the grammar.

The pairwise disjointness test is as follows:

For each nonterminal, \(A\), in the grammar that has more than one RHS, for each pair of rules, \(A \rightarrow \alpha_i\) and \(A \rightarrow \alpha_j\), it must be true that

\[
\text{FIRST}(\alpha_i) \cap \text{FIRST}(\alpha_j) = \emptyset
\]

(The intersection of the two sets, FIRST(\(\alpha_i\)) and FIRST(\(\alpha_j\)), must be empty.)

In other words, if a nonterminal \(A\) has more than one RHS, the first terminal symbol that can be generated in a derivation for each of them must be unique to that RHS. Consider the following rules:

\[
\begin{align*}
A & \rightarrow aB \mid bAb \mid Bb \\
B & \rightarrow cB \mid d
\end{align*}
\]

The FIRST sets for the RHSs of the A-rules are \{a\}, \{b\}, and \{c\}, \{d\}, which are clearly disjoint. Therefore, these rules pass the pairwise disjointness test. What this means, in terms of a recursive-descent parser, is that the code of the subprogram for parsing the nonterminal A can choose which RHS it is dealing with by seeing only the first terminal symbol of input (token) that is generated by the nonterminal. Now consider the rules

\[
\begin{align*}
A & \rightarrow aB \mid BAb \\
B & \rightarrow aB \mid b
\end{align*}
\]
The FIRST sets for the RHSs in the A-rules are \{a\} and \{a\}, \{b\} which are clearly not disjoint. So, these rules fail the pairwise disjointness test. In terms of the parser, the subprogram for A could not determine which RHS was being parsed by looking at the next symbol of input, because if it were an a, it could be either RHS. This issue is of course more complex if one or more of the RHSs begin with nonterminals.

In many cases, a grammar that fails the pairwise disjointness test can be modified so that it will pass the test. For example, consider the rule

\[ \text{<variable>} \rightarrow \text{identifier} \mid \text{identifier} \ [\text{<expression>}] \]

This states that a <variable> is either an identifier or an identifier followed by an expression in brackets (a subscript). These rules clearly do not pass the pairwise disjointness test, because both RHSs begin with the same terminal, identifier. This problem can be alleviated through a process called **left factoring**.

We now take an informal look at left factoring. Consider our rules for <variable>. Both RHSs begin with identifier. The parts that follow identifier in the two RHSs are \(\varepsilon\) (the empty string) and \([\text{<expression>}]\). The two rules can be replaced by the following two rules:

\[ \text{<variable>} \rightarrow \text{identifier} \ \text{<new>} \]

\[ \text{<new>} \rightarrow \varepsilon \mid [\text{<expression>}] \]

It is not difficult to see that together, these two rules generate the same language as the two rules with which we began. However, these two pass the pairwise disjointness test.

If the grammar is being used as the basis for a recursive-descent parser, an alternative to left factoring is available. With an EBNF extension, the problem disappears in a way that is very similar to the left factoring solution. Consider the original rules above for <variable>. The subscript can be made optional by placing it in square brackets, as in

\[ \text{<variable>} \rightarrow \text{identifier} \ [\ [\text{<expression}>] \ ] \]

In this rule, the outer brackets are metasymbols that indicate that what is inside is optional. The inner brackets are terminal symbols of the programming language being described. The point is that we replaced two rules with a single rule that generates the same language but passes the pairwise disjointness test.

A formal algorithm for left factoring can be found in Aho et al. (2006). Left factoring cannot solve all pairwise disjointness problems of grammars. In some cases, rules must be rewritten in other ways to eliminate the problem.

---

### 4.5 Bottom-Up Parsing

This section introduces the general process of bottom-up parsing and includes a description of the LR parsing algorithm.
4.5.1 The Parsing Problem for Bottom-Up Parsers

Consider the following grammar for arithmetic expressions:

\[ E \rightarrow E + T | T \]
\[ T \rightarrow T * F | F \]
\[ F \rightarrow (E) | \text{id} \]

Notice that this grammar generates the same arithmetic expressions as the example in Section 4.4. The difference is that this grammar is left recursive, which is acceptable to bottom-up parsers. Also note that grammars for bottom-up parsers normally do not include metasymbols such as those used to specify extensions to BNF. The following rightmost derivation illustrates this grammar:

\[ E \rightarrow E + T \]
\[ \rightarrow E + T * F \]
\[ \rightarrow E + T * \text{id} \]
\[ \rightarrow E + F * \text{id} \]
\[ \rightarrow E + \text{id} * \text{id} \]
\[ \rightarrow T + \text{id} * \text{id} \]
\[ \rightarrow F + \text{id} * \text{id} \]
\[ \rightarrow \text{id} + \text{id} * \text{id} \]

The underlined part of each sentential form in this derivation is the RHS that is rewritten as its corresponding LHS to get the previous sentential form. The process of bottom-up parsing produces the reverse of a rightmost derivation. So, in the example derivation, a bottom-up parser starts with the last sentential form (the input sentence) and produces the sequence of sentential forms from there until all that remains is the start symbol, which in this grammar is E. In each step, the task of the bottom-up parser is to find the specific RHS, the handle, in the sentential form that must be rewritten to get the next (previous) sentential form. As mentioned earlier, a right sentential form may include more than one RHS. For example, the right sentential form

\[ E + T * \text{id} \]

includes three RHSs, \( E + T \), \( T \), and \( \text{id} \). Only one of these is the handle. For example, if the RHS \( E + T \) were chosen to be rewritten in this sentential form, the resulting sentential form would be \( E * \text{id} \), but \( E * \text{id} \) is not a legal right sentential form for the given grammar.

The handle of a right sentential form is unique. The task of a bottom-up parser is to find the handle of any given right sentential form that can be generated by its associated grammar. Formally, handle is defined as follows:

**Definition:** \( \beta \) is the **handle** of the right sentential form \( \gamma = \alpha \beta w \) if and only if \( S \Rightarrow *_{rm} \alpha A w \Rightarrow_{m} \alpha \beta w \)

In this definition, \( \Rightarrow_{m} \) specifies a rightmost derivation step, and \( \Rightarrow *_{rm} \) specifies zero or more rightmost derivation steps. Although the definition of a
handle is mathematically concise, it provides little help in finding the handle of a given right sentential form. In the following, we provide the definitions of several substrings of sentential forms that are related to handles. The purpose of these is to provide some intuition about handles.

Definition: $\beta$ is a phrase of the right sentential form $\gamma$ if and only if
$S \Rightarrow^\ast \gamma = \alpha_1 A \alpha_2 \Rightarrow^+ \alpha_1 \beta \alpha_2$

In this definition, $\Rightarrow^+ \alpha$ means one or more derivation steps.

Definition: $\beta$ is a simple phrase of the right sentential form $\gamma$ if and only if
$S \Rightarrow^\ast \gamma = \alpha_1 A \alpha_2 \Rightarrow \alpha_1 \beta \alpha_2$

If these two definitions are compared carefully, it is clear that they differ only in the last derivation specification. The definition of phrase uses one or more steps, while the definition of simple phrase uses exactly one step.

The definitions of phrase and simple phrase may appear to have the same lack of practical value as that of a handle, but that is not true. Consider what a phrase is relative to a parse tree. It is the string of all of the leaves of the partial parse tree that is rooted at one particular internal node of the whole parse tree. A simple phrase is just a phrase that takes a single derivation step from its root nonterminal node. In terms of a parse tree, a phrase can be derived from a single nonterminal in one or more tree levels, but a simple phrase can be derived in just a single tree level. Consider the parse tree shown in Figure 4.3.

Figure 4.3
A parse tree for $E + T \ast id$

The leaves of the parse tree in Figure 4.3 comprise the sentential form $E + T \ast id$. Because there are three internal nodes, there are three phrases. Each internal node is the root of a subtree, whose leaves are a phrase. The root node of the whole parse tree, $E$, generates all of the resulting sentential form, $E + T \ast id$, which is a phrase. The internal node, $T$, generates the leaves $T \ast id$, which is another phrase. Finally, the internal node, $F$, generates $id$, which is also a phrase. So, the phrases of the sentential form $E + T \ast id$ are $E + T \ast id$, $T \ast id$, and $id$. Notice that phrases are not necessarily RHSs in the underlying grammar.
The simple phrases are a subset of the phrases. In the previous example, the only simple phrase is id. A simple phrase is always a RHS in the grammar.

The reason for discussing phrases and simple phrases is this: The handle of any rightmost sentential form is its leftmost simple phrase. So now we have a highly intuitive way to find the handle of any right sentential form, assuming we have the grammar and can draw a parse tree. This approach to finding handles is of course not practical for a parser. (If you already have a parse tree, why do you need a parser?) Its only purpose is to provide the reader with some intuitive feel for what a handle is, relative to a parse tree, which is easier than trying to think about handles in terms of sentential forms.

We can now consider bottom-up parsing in terms of parse trees, although the purpose of a parser is to produce a parse tree. Given the parse tree for an entire sentence, you easily can find the handle, which is the first thing to rewrite in the sentence to get the previous sentential form. Then the handle can be pruned from the parse tree and the process repeated. Continuing to the root of the parse tree, the entire rightmost derivation can be constructed.

### 4.5.2 Shift-Reduce Algorithms

Bottom-up parsers are often called shift-reduce algorithms, because shift and reduce are the two most common actions they specify. An integral part of every bottom-up parser is a stack. As with other parsers, the input to a bottom-up parser is the stream of tokens of a program and the output is a sequence of grammar rules. The shift action moves the next input token onto the parser’s stack. A reduce action replaces an RHS (the handle) on top of the parser’s stack by its corresponding LHS. Every parser for a programming language is a pushdown automaton (PDA), because a PDA is a recognizer for a context-free language. You need not be intimate with PDAs to understand how a bottom-up parser works, although it helps. A PDA is a very simple mathematical machine that scans strings of symbols from left to right. A PDA is so named because it uses a pushdown stack as its memory. PDAs can be used as recognizers for context-free languages. Given a string of symbols over the alphabet of a context-free language, a PDA that is designed for the purpose can determine whether the string is or is not a sentence in the language. In the process, the PDA can produce the information needed to construct a parse tree for the sentence.

With a PDA, the input string is examined, one symbol at a time, left to right. The input is treated very much as if it were stored in another stack, because the PDA never sees more than the leftmost symbol of the input.

Note that a recursive-descent parser is also a PDA. In that case, the stack is that of the run-time system, which records subprogram calls (among other things), which correspond to the nonterminals of the grammar.

### 4.5.3 LR Parsers

Many different bottom-up parsing algorithms have been devised. Most of them are variations of a process called LR. LR parsers use a relatively small program and a parsing table that is built for a specific programming language. The
original LR algorithm was designed by Donald Knuth (Knuth, 1965). This algorithm, which is sometimes called canonical LR, was not used in the years immediately following its publication because producing the required parsing table required large amounts of computer time and memory. Subsequently, several variations on the canonical LR table construction process were developed (DeRemer, 1971; DeRemer and Pennello, 1982). These are characterized by two properties: (1) They require far less computer resources to produce the required parsing table than the canonical LR algorithm, and (2) they work on smaller classes of grammars than the canonical LR algorithm.

There are three advantages to LR parsers:

1. They can be built for all programming languages.
2. They can detect syntax errors as soon as it is possible in a left-to-right scan.
3. The LR class of grammars is a proper superset of the class parsable by LL parsers (for example, many left recursive grammars are LR, but none are LL).

The only disadvantage of LR parsing is that it is difficult to produce by hand the parsing table for a given grammar for a complete programming language. This is not a serious disadvantage, however, for there are several programs available that take a grammar as input and produce the parsing table, as discussed later in this section.

Prior to the appearance of the LR parsing algorithm, there were a number of parsing algorithms that found handles of right sentential forms by looking both to the left and to the right of the substring of the sentential form that was suspected of being the handle. Knuth’s insight was that one could effectively look to the left of the suspected handle all the way to the bottom of the parse stack to determine whether it was the handle. But all of the information in the parse stack that was relevant to the parsing process could be represented by a single state, which could be stored on the top of the stack. In other words, Knuth discovered that regardless of the length of the input string, the length of the sentential form, or the depth of the parse stack, there were only a relatively small number of different situations, as far as the parsing process is concerned. Each situation could be represented by a state and stored in the parse stack, one state symbol for each grammar symbol on the stack. At the top of the stack would always be a state symbol, which represented the relevant information from the entire history of the parse, up to the current time. We will use subscripted uppercase S’s to represent the parser states.

Figure 4.4 shows the structure of an LR parser. The contents of the parse stack for an LR parser have the following form:

\[S_0X_1S_1X_2 \ldots X_mS_m\] (top)

where the S’s are state symbols and the X’s are grammar symbols. An LR parser configuration is a pair of strings (stack, input), with the detailed form

\[(S_0X_1S_1X_2S_2 \ldots X_mS_m, a_ia_{i+1} \ldots a_n$)\]
Notice that the input string has a dollar sign at its right end. This sign is put there during initialization of the parser. It is used for normal termination of the parser. Using this parser configuration, we can formally define the LR parser process, which is based on the parsing table.

An LR parsing table has two parts, named ACTION and GOTO. The ACTION part of the table specifies most of what the parser does. It has state symbols as its row labels and the terminal symbols of the grammar as its column labels. Given a current parser state, which is represented by the state symbol on top of the parse stack, and the next symbol (token) of input, the parse table specifies what the parser should do. The two primary parser actions are shift and reduce. Either the parser shifts the next input symbol onto the parse stack or it already has the handle on top of the stack, which it reduces to the LHS of the rule whose RHS is the same as the handle. Two other actions are possible: accept, which means the parser has successfully completed the parse of the input, and error, which means the parser has detected a syntax error.

The rows of the GOTO part of the LR parsing table have state symbols as labels. This part of the table has nonterminals as column labels. The values in the GOTO part of the table indicate which state symbol should be pushed onto the parse stack after a reduction has been completed, which means the handle has been removed from the parse stack and the new nonterminal has been pushed onto the parse stack. The specific symbol is found at the row whose label is the state symbol on top of the parse stack after the handle and its associated state symbols have been removed. The column of the GOTO table that is used is the one with the label, that is the LHS of the rule used in the reduction.

Consider the traditional grammar for arithmetic expressions that follows:

1. $E \rightarrow E + T$
2. $E \rightarrow T$
3. $T \rightarrow T \ast F$
4. $T \rightarrow F$
5. $F \rightarrow (E)$
6. $F \rightarrow id$
4.5 Bottom-Up Parsing

The rules of this grammar are numbered to provide a simple way to reference them in a parsing table.

Figure 4.5 shows the LR parsing table for this grammar. Abbreviations are used for the actions: \( R \) for reduce and \( S \) for shift. \( R4 \) means reduce using rule 4; \( S6 \) means shift the next symbol of input onto the stack and push state \( S6 \) onto the stack. Empty positions in the ACTION table indicate syntax errors. In a complete parser, these could have calls to error-handling routines.

LR parsing tables can easily be constructed using a software tool, such as yacc (Johnson, 1975), which takes the grammar as input. Although LR parsing tables can be produced by hand, for a grammar of a real programming language, the task would be lengthy, tedious, and error prone. For real compilers, LR parsing tables are always generated with software tools.

The initial configuration of an LR parser is

\[(S_0, a_1 \ldots a_n$)\]

The parser actions are informally defined as follows:

1. The Shift process is simple: The next symbol of input is pushed onto the stack, along with the state symbol that is part of the Shift specification in the ACTION table.

2. For a Reduce action, the handle must be removed from the stack. Because for every grammar symbol on the stack there is a state symbol, the number of symbols removed from the stack is twice the number

---

![Figure 4.5](image)

The LR parsing table for an arithmetic expression grammar.
of symbols in the handle. After removing the handle and its associated state symbols, the LHS of the rule is pushed onto the stack. Finally, the GOTO table is used, with the row label being the symbol that was exposed when the handle and its state symbols were removed from the stack, and the column label being the nonterminal that is the LHS of the rule used in the reduction.

3. When the action is Accept, the parse is complete and no errors were found.

4. When the action is Error, the parser calls an error-handling routine.

Although there are many parsing algorithms based on the LR concept, they differ only in the construction of the parsing table. All LR parsers use this same parsing algorithm.

Perhaps the best way to become familiar with the LR parsing process is through an example. Initially, the parse stack has the single symbol 0, which represents state 0 of the parser. The input contains the input string with an end marker, in this case a dollar sign, attached to its right end. At each step, the parser actions are dictated by the top (rightmost in Figure 4.4) symbol of the parse stack and the next (leftmost in Figure 4.4) token of input. The correct action is chosen from the corresponding cell of the ACTION part of the parse table. The GOTO part of the parse table is used after a reduction action. Recall that GOTO is used to determine which state symbol is placed on the parse stack after a reduction.

Following is a trace of a parse of the string id + id * id, using the LR parsing algorithm and the parsing table shown in Figure 4.5.

<table>
<thead>
<tr>
<th>Stack</th>
<th>Input</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>id + id * id $</td>
<td>Shift 5</td>
</tr>
<tr>
<td>0id5</td>
<td>+ id * id $</td>
<td>Reduce 6 (use GOTO[0, F])</td>
</tr>
<tr>
<td>0F3</td>
<td>+ id * id $</td>
<td>Reduce 4 (use GOTO[0, T])</td>
</tr>
<tr>
<td>0T2</td>
<td>+ id * id $</td>
<td>Reduce 2 (use GOTO[0, E])</td>
</tr>
<tr>
<td>0E1</td>
<td>+ id * id $</td>
<td>Shift 6</td>
</tr>
<tr>
<td>0E1+6</td>
<td>id * id $</td>
<td>Shift 5</td>
</tr>
<tr>
<td>0E1+6id5</td>
<td>* id $</td>
<td>Reduce 6 (use GOTO[6, F])</td>
</tr>
<tr>
<td>0E1+6F3</td>
<td>* id $</td>
<td>Reduce 4 (use GOTO[6, T])</td>
</tr>
<tr>
<td>0E1+6T9</td>
<td>* id $</td>
<td>Shift 7</td>
</tr>
<tr>
<td>0E1+6T9*7</td>
<td>id $</td>
<td>Shift 5</td>
</tr>
<tr>
<td>0E1+6T9*7id5</td>
<td>$</td>
<td>Reduce 6 (use GOTO[7, F])</td>
</tr>
<tr>
<td>0E1+6T9*7F10</td>
<td>$</td>
<td>Reduce 3 (use GOTO[6, T])</td>
</tr>
<tr>
<td>0E1+6T9</td>
<td>$</td>
<td>Reduce 1 (use GOTO[0, E])</td>
</tr>
<tr>
<td>0E1</td>
<td>$</td>
<td>Accept</td>
</tr>
</tbody>
</table>

The algorithms to generate LR parsing tables from given grammars, which are described in Aho et al. (2006), are not overly complex but are beyond the scope of a book on programming languages. As stated previously, there are a number of different software systems available to generate LR parsing tables.
Syntax analysis is a common part of language implementation, regardless of the implementation approach used. Syntax analysis is normally based on a formal syntax description of the language being implemented. A context-free grammar, which is also called BNF, is the most common approach for describing syntax. The task of syntax analysis is usually divided into two parts: lexical analysis and syntax analysis. There are several reasons for separating lexical analysis—namely, simplicity, efficiency, and portability.

A lexical analyzer is a pattern matcher that isolates the small-scale parts of a program, which are called lexemes. Lexemes occur in categories, such as integer literals and names. These categories are called tokens. Each token is assigned a numeric code, which along with the lexeme is what the lexical analyzer produces. There are three distinct approaches to constructing a lexical analyzer: using a software tool to generate a table for a table-driven analyzer, building such a table by hand, and writing code to implement a state diagram description of the tokens of the language being implemented. The state diagram for tokens can be reasonably small if character classes are used for transitions, rather than having transitions for every possible character from every state node. Also, the state diagram can be simplified by using a table lookup to recognize reserved words.

Syntax analyzers have two goals: to detect syntax errors in a given program and to produce a parse tree, or possibly only the information required to build such a tree, for a given program. Syntax analyzers are either top-down, meaning they construct leftmost derivations and a parse tree in top-down order, or bottom-up, in which case they construct the reverse of a rightmost derivation and a parse tree in bottom-up order. Parsers that work for all unambiguous grammars have complexity $O(n^3)$. However, parsers used for implementing syntax analyzers for programming languages work on subclasses of unambiguous grammars and have complexity $O(n)$.

A recursive-descent parser is an LL parser that is implemented by writing code directly from the grammar of the source language. EBNF is ideal as the basis for recursive-descent parsers. A recursive-descent parser has a subprogram for each nonterminal in the grammar. The code for a given grammar rule is simple if the rule has a single RHS. The RHS is examined left to right. For each nonterminal, the code calls the associated subprogram for that nonterminal, which parses whatever the nonterminal generates. For each terminal, the code compares the terminal with the next token of input. If they match, the code simply calls the lexical analyzer to get the next token. If they do not, the subprogram reports a syntax error. If a rule has more than one RHS, the subprogram must first determine which RHS it should parse. It must be possible to make this determination on the basis of the next token of input.

Two distinct grammar characteristics prevent the construction of a recursive-descent parser based on the grammar. One of these is left recursion. The process of eliminating direct left recursion from a grammar is relatively simple. Although we do not cover it, an algorithm exists to remove both direct
and indirect left recursion from a grammar. The other problem is detected with the pairwise disjointness test, which tests whether a parsing subprogram can determine which RHS is being parsed on the basis of the next token of input. Some grammars that fail the pairwise disjointness test often can be modified to pass it, using left factoring.

The parsing problem for bottom-up parsers is to find the substring of the current sentential form that must be reduced to its associated LHS to get the next (previous) sentential form in the rightmost derivation. This substring is called the handle of the sentential form. A parse tree can provide an intuitive basis for recognizing a handle. A bottom-up parser is a shift-reduce algorithm, because in most cases it either shifts the next lexeme of input onto the parse stack or reduces the handle that is on top of the stack.

The LR family of shift-reduce parsers is the most commonly used bottom-up parsing approach for programming languages, because parsers in this family have several advantages over alternatives. An LR parser uses a parse stack, which contains grammar symbols and state symbols to maintain the state of the parser. The top symbol on the parse stack is always a state symbol that represents all of the information in the parse stack that is relevant to the parsing process. LR parsers use two parsing tables: ACTION and GOTO. The ACTION part specifies what the parser should do, given the state symbol on top of the parse stack and the next token of input. The GOTO table is used to determine which state symbol should be placed on the parse stack after a reduction has been done.

**REVIEW QUESTIONS**

1. What are the reasons why using BNF is advantageous over using an informal syntax description?
2. How does a lexical analyzer serve as the front end of a syntax analyzer?
3. Define finite automata and regular grammar.
4. How can you construct a lexical analyzer with a state diagram?
5. Describe briefly the three approaches to building a lexical analyzer.
6. What are the different grammar symbols for formal languages?
7. Why are character classes used, rather than individual characters, for the letter and digit transitions of a state diagram for a lexical analyzer?
8. What are the two distinct goals of syntax analysis?
9. Describe the complexity of parsing algorithms.
10. Describe the recursive-descent parser.
11. What do the two Ls in LL algorithm specify?
12. State and explain the convention followed for writing a recursive-descent parsing subprogram.
13. Why are named constants used, rather than numbers, for token codes?
14. Describe how a recursive-descent parsing subprogram is written for a rule with a single RHS.
15. Explain the two grammar characteristics that prohibit them from being used as the basis for a top-down parser.
16. What is direct left recursion?
17. Describe the pairwise disjointness test.
18. What is the limitation of left factoring?
19. What is a phrase of a right sentential form?
20. What is the difference between a simple phrase and a phrase of the right sentential form?
21. What is the feature of the handle of a right sentential form?
22. What is the mathematical machine on which both top-down and bottom-up parsers are based?
23. What is the disadvantage of an LR parser?
24. Why is a bottom-up parser often called a shift-reduce algorithm?
25. Describe the purpose of a parse stack in an LR parser.
26. Describe the properties of the variations on the canonical LR table construction process.
27. Why is every parser for a programming language a pushdown automaton?

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**Problem Set**

1. Perform the pairwise disjointness test for the following grammar rules.
   a. \(A \rightarrow aB \mid b \mid cBB\)
   b. \(B \rightarrow aB \mid bA \mid aBb\)
   c. \(C \rightarrow aaA \mid b \mid caB\)
2. Perform the pairwise disjointness test for the following grammar rules.
   a. \(S \rightarrow aSb \mid bAA\)
   b. \(A \rightarrow b[aB] \mid a\)
   c. \(B \rightarrow aB \mid a\)
3. Show a trace of the recursive descent parser given in Section 4.4.1 for the string \(a + b * c\).
4. Show a trace of the recursive descent parser given in Section 4.4.1 for the string \(a * (b + c)\).
5. Given the following grammar and the right sentential form, draw a parse tree and show the phrases and simple phrases, as well as the handle.
   \(S \rightarrow aAb \mid bBA\)
   \(A \rightarrow ab \mid aAB\)
   \(B \rightarrow aB \mid b\)
   a. \(aaAbBb\)
   b. \(bBab\)
   c. \(aaAbBb\)
6. Given the following grammar and the right sentential form, draw a parse tree and show the phrases and simple phrases, as well as the handle.

\[
S \rightarrow AbB \mid bAc \quad A \rightarrow Ab \quad aBB \quad B \rightarrow Ac \quad cBb \mid c \\
\]

a. \( aAccbcbc \)
b. \( AbcaBccb \)
c. \( baBcBbbc \)

7. Show a complete parse, including the parse stack contents, input string, and action for the string \( id \ast (id + id) \), using the grammar and parse table in Section 4.5.3.

8. Show a complete parse, including the parse stack contents, input string, and action for the string \( (id + id) \ast id \), using the grammar and parse table in Section 4.5.3.

9. Get the algorithm to remove the indirect left recursion from a grammar from Aho et al. (2006). Use this algorithm to remove all left recursion from the following grammar:

\[
S \rightarrow Aa \mid Bb \\
A \rightarrow Abc \mid c \\
B \rightarrow bb
\]

**Programming Exercises**

1. Design a state diagram to recognize one form of the comments of the C-based programming languages, those that begin with /* and end with */.

2. Design a state diagram to recognize the floating-point literals of your favorite programming language.

3. Write and test the code to implement the state diagram of Problem 1.

4. Write and test the code to implement the state diagram of Problem 2.

5. Modify the lexical analyzer given in Section 4.2 to recognize the following list of reserved words and return their respective token codes:

\[
for (FOR_CODE, 30), \text{if} (IF_CODE, 31), \text{else} (ELSE_CODE, 32), \text{while} (WHILE_CODE, 33), \text{do} (DO_CODE, 34), \text{int} (INT_CODE, 35), \text{float} (FLOAT_CODE, 36), \text{switch} (SWITCH_CODE, 37).
\]

6. Convert the lexical analyzer (which is written in C) given in Section 4.2 to Java.

7. Convert the recursive descent parser routines for <expr>, <term>, and <factor> given in Section 4.4.1 to Java.

8. For those rules that pass the test in Problem 1, write a recursive-descent parsing subprogram that parses the language generated by the rules. Assume you have a lexical analyzer named lex and an error-handling subprogram named error, which is called whenever a syntax error is detected.

9. For those rules that pass the test in Problem 2, write a recursive-descent parsing subprogram that parses the language generated by the rules.
Assume you have a lexical analyzer named `lex` and an error-handling subprogram named `error`, which is called whenever a syntax error is detected.

10. Implement and test the LR parsing algorithm given in Section 4.5.3.
11. Write an EBNF rule that describes the `while` statement of Java or C++. Write the recursive-descent subprogram in Java or C++ for this rule.
12. Write an EBNF rule that describes the `for` statement of Java or C++. Write the recursive-descent subprogram in Java or C++ for this rule.
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5.1 Introduction
5.2 Names
5.3 Variables
5.4 The Concept of Binding
5.5 Scope
5.6 Scope and Lifetime
5.7 Referencing Environments
5.8 Named Constants
This chapter introduces the fundamental semantic issues of variables. It begins by describing the nature of names and special words in programming languages. The attributes of variables, including type, address, and value, are then discussed, including the issue of aliases. The important concepts of binding and binding times are introduced next, including the different possible binding times for variable attributes and how they define four different categories of variables. Following that, two very different scoping rules for names, static and dynamic, are described, along with the concept of a referencing environment of a statement. Finally, named constants and variable initialization are discussed.

5.1 Introduction

Imperative programming languages are, to varying degrees, abstractions of the underlying von Neumann computer architecture. The architecture’s two primary components are its memory, which stores both instructions and data, and its processor, which provides operations for modifying the contents of the memory. The abstractions in a language for the memory cells of the machine are variables. In some cases, the characteristics of the abstractions are very close to the characteristics of the cells; an example of this is an integer variable, which is usually represented directly in one or more bytes of memory. In other cases, the abstractions are far removed from the organization of the hardware memory, as with a three-dimensional array, which requires a software mapping function to support the abstraction.

A variable can be characterized by a collection of properties, or attributes, the most important of which is type, a fundamental concept in programming languages. Designing the data types of a language requires that a variety of issues be considered. (Data types are discussed in Chapter 6.) Among the most important of these issues are the scope and lifetime of variables.

Functional programming languages allow expressions to be named. These named expressions appear like assignments to variable names in imperative languages, but are fundamentally different in that they cannot be changed. So, they are like the named constants of the imperative languages. Pure functional languages do not have variables that are like those of the imperative languages. However, many functional languages do include such variables.

In the remainder of this book, families of languages will often be referred to as if they were single languages. For example, Fortran will mean all of the versions of Fortran. This is also the case for Ada. Likewise, a reference to C will mean the original version of C, as well as C89 and C99. When a specific version of a language is named, it is because it is different from the other family members within the topic being discussed. If we add a plus sign (+) to the name of a version of a language, we mean all versions of the language beginning with the one named. For example, Fortran 95+ means all versions of Fortran beginning with Fortran 95. The phrase C-based languages will be used to refer to C, Objective-C, C++, Java, and C#.

1. We were tempted to include the scripting languages JavaScript and PHP as C-based languages, but decided they were just a bit too different from their ancestors.
Before beginning our discussion of variables, the design of one of the fundamental attributes of variables, names, must be covered. Names are also associated with subprograms, formal parameters, and other program constructs. The term *identifier* is often used interchangeably with *name*.

### 5.2.1 Design Issues

The following are the primary design issues for names:

- Are names case sensitive?
- Are the special words of the language reserved words or keywords?

These issues are discussed in the following two subsections, which also include examples of several design choices.

### 5.2.2 Name Forms

A *name* is a string of characters used to identify some entity in a program. C99 has no length limitation on its internal names, but only the first 63 are significant. External names in C99 (those defined outside functions, which must be handled by the linker) are restricted to 31 characters. Names in Java and C# have no length limit, and all characters in them are significant. C++ does not specify a length limit on names, although implementors sometimes do.

Names in most programming languages have the same form: a letter followed by a string consisting of letters, digits, and underscore characters (\_). Although the use of underscore characters to form names was widely used in the 1970s and 1980s, that practice is now far less popular. In the C-based languages, it has to a large extent been replaced by the so-called *camel notation*, in which all of the words of a multiple-word name except the first are capitalized, as in *myStack*.

Note that the use of underscores and mixed case in names is a programming style issue, not a language design issue.

All variable names in PHP must begin with a dollar sign. In Perl, the special character at the beginning of a variable’s name, $, @, or %, specifies its type (although in a different sense than in other languages). In Ruby, special characters at the beginning of a variable’s name, @ or @@, indicate that the variable is an instance or a class variable, respectively.

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2. It is called “camel” because words written in it often have embedded uppercase letters, which look like a camel’s humps.
In many languages, notably the C-based languages, uppercase and lowercase letters in names are distinct; that is, names in these languages are **case sensitive**. For example, the following three names are distinct in C++: *rose*, *ROSE*, and *Rose*. To some people, this is a serious detriment to readability, because names that look very similar in fact denote different entities. In that sense, case sensitivity violates the design principle that language constructs that look similar should have similar meanings. But in languages whose variable names are case-sensitive, although *Rose* and *rose* look similar, there is no connection between them.

Obviously, not everyone agrees that case sensitivity is bad for names. In C, the problems of case sensitivity are avoided by the convention that variable names do not include uppercase letters. In Java and C#, however, the problem cannot be escaped because many of the predefined names include both uppercase and lowercase letters. For example, the Java method for converting a string to an integer value is `parseInt`, and spellings such as `ParseInt` and `parseint` are not recognized. This is a problem of writability rather than readability, because the need to remember specific case usage makes it more difficult to write correct programs. It is a kind of intolerance on the part of the language designer, which is enforced by the compiler.

### 5.2.3 Special Words

Special words in programming languages are used to make programs more readable by naming actions to be performed. They also are used to separate the syntactic parts of statements and programs. In most languages, special words are classified as reserved words, which means they cannot be redefined by programmers. But in some, such as Fortran, they are only keywords, which means they can be redefined.

A **reserved word** is a special word of a programming language that cannot be used as a name. There is one potential problem with reserved words: If the language includes a large number of reserved words, the user may have difficulty making up names that are not reserved. The best example of this is COBOL, which has 300 reserved words. Unfortunately, some of the most commonly chosen names by programmers are in the list of reserved words—for example, `LENGTH`, `BOTTOM`, `DESTINATION`, and `COUNT`.

In program code examples in this book, reserved words are presented in **boldface**.

In most languages, names that are defined in other program units, such as Java packages and C and C++ libraries, can be made visible to a program. These names are predefined, but visible only if explicitly imported. Once imported, they cannot be redefined.

### 5.3 Variables

A program variable is an abstraction of a computer memory cell or collection of cells. Programmers often think of variables as names for memory locations, but there is much more to a variable than just a name.
The move from machine languages to assembly languages was largely one of replacing absolute numeric memory addresses for data with names, making programs far more readable and therefore easier to write and maintain. Assembly languages also provided an escape from the problem of manual absolute addressing, because the translator that converted the names to actual addresses also chose those addresses.

A variable can be characterized as a sextuple of attributes: (name, address, value, type, lifetime, and scope). Although this may seem too complicated for such an apparently simple concept, it provides the clearest way to explain the various aspects of variables.

Our discussion of variable attributes will lead to examinations of the important related concepts of aliases, binding, binding times, declarations, scoping rules, and referencing environments.

The name, address, type, and value attributes of variables are discussed in the following subsections. The lifetime and scope attributes are discussed in Sections 5.4.3 and 5.5, respectively.

### 5.3.1 Name

Variable names are the most common names in programs. They were discussed at length in Section 5.2 in the general context of entity names in programs. Most variables have names. The ones that do not are discussed in Section 5.4.3.3.

### 5.3.2 Address

The address of a variable is the machine memory address with which it is associated. This association is not as simple as it may at first appear. In many languages, it is possible for the same variable to be associated with different addresses at different times during the execution of the program. For example, if a subprogram has a local variable that is allocated from the run-time stack when the subprogram is called, different calls may result in that variable having different addresses. These are in a sense different instantiations of the same variable.

The process of associating variables with addresses is further discussed in Section 5.4.3. An implementation model for subprograms and their activations is discussed in Chapter 10.

The address of a variable is sometimes called its l-value, because the address is what is required when the name of a variable appears in the left side of an assignment.

It is possible to have multiple variables that have the same address. When more than one variable name can be used to access the same memory location, the variables are called aliases. Aliasing is a hindrance to readability because it allows a variable to have its value changed by an assignment to a different variable. For example, if variables named total and sum are aliases, any change to the value of total also changes the value of sum and vice versa. A reader of the program must always remember that total and sum are different names.
for the same memory cell. Because there can be any number of aliases in a program, this may be very difficult in practice. Aliasing also makes program verification more difficult.

Aliases can be created in programs in several different ways. One common way in C and C++ is with their union types. Unions are discussed at length in Chapter 6.

Two pointer variables are aliases when they point to the same memory location. The same is true for reference variables. This kind of aliasing is simply a side effect of the nature of pointers and references. When a C++ pointer is set to point at a named variable, the pointer, when dereferenced, and the variable’s name are aliases.

Aliasing can be created in many languages through subprogram parameters. These kinds of aliases are discussed in Chapter 9.

The time when a variable becomes associated with an address is very important to an understanding of programming languages. This subject is discussed in Section 5.4.3.

5.3.3 Type

The type of a variable determines the range of values the variable can store and the set of operations that are defined for values of the type. For example, the int type in Java specifies a value range of −2,147,483,648 to 2,147,483,647 and arithmetic operations for addition, subtraction, multiplication, division, and modulus.

5.3.4 Value

The value of a variable is the contents of the memory cell or cells associated with the variable. It is convenient to think of computer memory in terms of abstract cells, rather than physical cells. The physical cells, or individually addressable units, of most contemporary computer memories are byte size, with a byte having eight bits. This size is too small for most program variables. An abstract memory cell has the size required by the variable with which it is associated. For example, although floating-point values may occupy four physical bytes in a particular implementation of a particular language, a floating-point value is thought of as occupying a single abstract memory cell. The value of each simple nonstructured type is considered to occupy a single abstract cell. Henceforth, the term memory cell will mean abstract memory cell.

A variable’s value is sometimes called its r-value because it is what is required when the name of the variable appears in the right side of an assignment statement. To access the r-value, the l-value must be determined first. Such determinations are not always simple. For example, scoping rules can greatly complicate matters, as is discussed in Section 5.5.
5.4 The Concept of Binding

A **binding** is an association between an attribute and an entity, such as between a variable and its type or value, or between an operation and a symbol. The time at which a binding takes place is called **binding time**. Binding and binding times are prominent concepts in the semantics of programming languages. Bindings can take place at language design time, language implementation time, compile time, load time, link time, or run time. For example, the asterisk symbol (*) is usually bound to the multiplication operation at language design time. A data type, such as `int` in C, is bound to a range of possible values at language implementation time. At compile time, a variable in a Java program is bound to a particular data type. A variable may be bound to a storage cell when the program is loaded into memory. That same binding does not happen until run time in some cases, as with variables declared in Java methods. A call to a library subprogram is bound to the subprogram code at link time.

Consider the following C++ assignment statement:

```cpp
count = count + 5;
```

Some of the bindings and their binding times for the parts of this assignment statement are as follows:

- The type of `count` is bound at compile time.
- The set of possible values of `count` is bound at compiler design time.
- The meaning of the operator symbol `+` is bound at compile time, when the types of its operands have been determined.
- The internal representation of the literal 5 is bound at compiler design time.
- The value of `count` is bound at execution time with this statement.

A complete understanding of the binding times for the attributes of program entities is a prerequisite for understanding the semantics of a programming language. For example, to understand what a subprogram does, one must understand how the actual parameters in a call are bound to the formal parameters in its definition. To determine the current value of a variable, it may be necessary to know when the variable was bound to storage and with which statement or statements.

### 5.4.1 Binding of Attributes to Variables

A binding is **static** if it first occurs before run time begins and remains unchanged throughout program execution. If the binding first occurs during run time or can change in the course of program execution, it is called **dynamic**. The physical binding of a variable to a storage cell in a virtual memory environment is complex, because the page or segment of the address space in which the cell resides may be moved in and out of memory many times during program execution. In a sense, such variables are bound and unbound repeatedly. These
bindings, however, are maintained by computer hardware, and the changes are invisible to the program and the user. Because they are not important to the discussion, we are not concerned with these hardware bindings. The essential point is to distinguish between static and dynamic bindings.

### 5.4.2 Type Bindings

Before a variable can be referenced in a program, it must be bound to a data type. The two important aspects of this binding are how the type is specified and when the binding takes place. Types can be specified statically through some form of explicit or implicit declaration.

#### 5.4.2.1 Static Type Binding

An **explicit declaration** is a statement in a program that lists variable names and specifies that they are a particular type. An **implicit declaration** is a means of associating variables with types through default conventions, rather than declaration statements. In this case, the first appearance of a variable name in a program constitutes its implicit declaration. Both explicit and implicit declarations create static bindings to types.

Most widely used programming languages that use static type binding exclusively and were designed since the mid-1960s require explicit declarations of all variables (Visual Basic and ML are two exceptions).

Implicit variable type binding is done by the language processor, either a compiler or an interpreter. There are several different bases for implicit variable type bindings. The simplest of these is naming conventions. In this case, the compiler or interpreter binds a variable to a type based on the syntactic form of the variable's name.

Although they are a minor convenience to programmers, implicit declarations can be detrimental to reliability because they prevent the compilation process from detecting some typographical and programmer errors.

Some of the problems with implicit declarations can be avoided by requiring names for specific types to begin with particular special characters. For example, in Perl any name that begins with `$_` is a scalar, which can store either a string or a numeric value. If a name begins with `@`, it is an array; if it begins with a `%`, it is a hash structure.\(^3\) This creates different namespaces for different type variables. In this scenario, the names `@apple` and `%apple` are unrelated, because each is from a different namespace. Furthermore, a program reader always knows the type of a variable when reading its name.

Another kind of implicit type declarations uses context. This is sometimes called **type inference**. In the simpler case, the context is the type of the value assigned to the variable in a declaration statement. For example, in C# a `var` declaration of a variable must include an initial value, whose type is taken as the type of the variable. Consider the following declarations:

---

3. Both arrays and hashes are considered types—both can store any scalar in their elements.
The types of `sum`, `total`, and `name` are `int`, `float`, and `string`, respectively. Keep in mind that these are statically typed variables—their types are fixed for the lifetime of the unit in which they are declared.

Visual Basic and the functional languages ML, Haskell, OCaml, and F# also use type inferencing. In these functional languages, the context of the appearance of a name is the basis for determining its type. This kind of type inferencing is discussed in detail in Chapter 15.

### 5.4.2.2 Dynamic Type Binding

With dynamic type binding, the type of a variable is not specified by a declaration statement, nor can it be determined by the spelling of its name. Instead, the variable is bound to a type when it is assigned a value in an assignment statement. When the assignment statement is executed, the variable being assigned is bound to the type of the value of the expression on the right side of the assignment. Such an assignment may also bind the variable to an address and a memory cell, because different type values may require different amounts of storage. Any variable can be assigned any type value. Furthermore, a variable’s type can change any number of times during program execution. It is important to realize that the type of a variable whose type is dynamically bound may be temporary.

When the type of a variable is statically bound, the name of the variable can be thought of being bound to a type, in the sense that the type and name of a variable are simultaneously bound. However, when a variable’s type is dynamically bound, its name can be thought of as being only temporarily bound to a type. In reality, the names of variables are never bound to types. Names can be bound to variables and variables can be bound to types.

Languages in which types are dynamically bound are dramatically different from those in which types are statically bound. The primary advantage of dynamic binding of variables to types is that it provides more programming flexibility. For example, a program to process numeric data in a language that uses dynamic type binding can be written as a generic program, meaning that it is capable of dealing with data of any numeric type. Whatever type data is input will be acceptable, because the variables in which the data are to be stored can be bound to the correct type when the data is assigned to the variables after input. By contrast, because of static binding of types, one cannot write a C program to process data without knowing the type of that data.

Before the mid-1990s, the most commonly used programming languages used static type binding, the primary exceptions being some functional languages such as Lisp. However, since then there has been a significant shift to languages that use dynamic type binding. In Python, Ruby, JavaScript, and PHP, type binding is dynamic. For example, a JavaScript script may contain the following statement:

```javascript
list = [10.2, 3.5];
```
Regardless of the previous type of the variable named `list`, this assignment causes it to become the name of a single-dimensioned array of length 2. If the statement

```csharp
list = 47;
```

followed the previous example assignment, `list` would become the name of a scalar variable.

The option of dynamic type binding was included in C# 2010. A variable can be declared to use dynamic type binding by including the `dynamic` reserved word in its declaration, as in the following example:

```csharp
dynamic any;
```

This is similar, although also different from declaring `any` to have type `object`. It is similar in that `any` can be assigned a value of any type, just as if it were declared `object`. It is different in that it is not useful for several different situations of interoperation; for example, with dynamically typed languages such as IronPython and IronRuby (.NET versions of Python and Ruby, respectively). However, it is useful when data of unknown type come into a program from an external source. Class members, properties, method parameters, method return values, and local variables can all be declared `dynamic`.

In pure object-oriented languages—for example, Ruby—all variables are references and do not have types; all data are objects and any variable can reference any object. Variables in such languages are, in a sense, all the same type—they are references. However, unlike the references in Java, which are restricted to referencing one specific type of value, variables in Ruby can reference any object.

There are two disadvantages to dynamic type binding. First, it causes programs to be less reliable, because the error-detection capability of the compiler is diminished relative to a compiler for a language with static type bindings. Dynamic type binding allows any variable to be assigned a value of any type. Incorrect types of right sides of assignments are not detected as errors; rather, the type of the left side is simply changed to the incorrect type. For example, suppose that in a particular JavaScript program, `i` and `x` are currently the names of scalar numeric variables and `y` is currently the name of an array. Furthermore, suppose that the program needs the assignment statement

```javascript
i = x;
```

but because of a keying error, it has the assignment statement

```javascript
i = y;
```

In JavaScript (or any other language that uses dynamic type binding), no error is detected in this statement by the interpreter—the type of the variable named `i` is simply changed to an array. But later uses of `i` will expect it to be a scalar, and correct results will be impossible. In a language with static type binding,
such as Java, the compiler would detect the error in the assignment $i = y$, and the program would not get to execution.

Note that this disadvantage is also present to some extent in some languages that use static type binding, such as C and C++, which in many cases automatically convert the type of the RHS of an assignment to the type of the LHS.

Perhaps the greatest disadvantage of dynamic type binding is cost. The cost of implementing dynamic attribute binding is considerable, particularly in execution time. Type checking must be done at run time. Furthermore, every variable must have a run-time descriptor associated with it to maintain the current type. The storage used for the value of a variable must be of varying size, because different type values require different amounts of storage.

Finally, languages that have dynamic type binding for variables are usually implemented using pure interpreters rather than compilers. Computers do not have instructions whose operand types are not known at compile time. Therefore, a compiler cannot build machine instructions for the expression $A + B$ if the types of $A$ and $B$ are not known at compile time. Pure interpretation typically takes at least 10 times as long as it does to execute equivalent machine code. Of course, if a language is implemented with a pure interpreter, the time to do dynamic type binding is hidden by the overall time of interpretation, so it seems less costly in that environment. On the other hand, languages with static type bindings are seldom implemented by pure interpretation, because programs in these languages can be easily translated to very efficient machine code versions.

### 5.4.3 Storage Bindings and Lifetime

The fundamental character of an imperative programming language is in large part determined by the design of the storage bindings for its variables. It is therefore important to have a clear understanding of these bindings.

The memory cell to which a variable is bound somehow must be taken from a pool of available memory. This process is called **allocation**. **Deallocation** is the process of placing a memory cell that has been unbound from a variable back into the pool of available memory.

The **lifetime** of a variable is the time during which the variable is bound to a specific memory location. So, the lifetime of a variable begins when it is bound to a specific cell and ends when it is unbound from that cell. To investigate storage bindings of variables, it is convenient to separate scalar (unstructured) variables into four categories, according to their lifetimes. These categories are named static, stack-dynamic, explicit heap-dynamic, and implicit heap-dynamic. In the following sections, we discuss the definitions of these four categories, along with their purposes, advantages, and disadvantages.

#### 5.4.3.1 Static Variables

**Static variables** are those that are bound to memory cells before program execution begins and remain bound to those same memory cells until program execution terminates. Statically bound variables have several valuable
applications in programming. Globally accessible variables are often used throughout the execution of a program, thus making it necessary to have them bound to the same storage during that execution. Sometimes it is convenient to have subprograms that are history sensitive. Such a subprogram must have local static variables.

One advantage of static variables is efficiency. All addressing of static variables can be direct; other kinds of variables often require indirect addressing, which is slower. Also, no run-time overhead is incurred for allocation and deallocation of static variables, although this time is often negligible.

One disadvantage of static binding to storage is reduced flexibility; in particular, a language that has only static variables cannot support recursive subprograms. Another disadvantage is that storage cannot be shared among variables. For example, suppose a program has two subprograms, both of which require large arrays. Furthermore, suppose that the two subprograms are never active at the same time. If the arrays are static, they cannot share the same storage for their arrays.

C and C++ allow programmers to include the `static` specifier on a variable definition in a function, making the variables it defines static. Note that when the `static` modifier appears in the declaration of a variable in a class definition in C++, Java, and C#, it also implies that the variable is a class variable, rather than an instance variable. Class variables are created statically some time before the class is first instantiated.

### 5.4.3.2 Stack-Dynamic Variables

**Stack-dynamic variables** are those whose storage bindings are created when their declaration statements are elaborated, but whose types are statically bound. **Elaboration** of such a declaration refers to the storage allocation and binding process indicated by the declaration, which takes place when execution reaches the code to which the declaration is attached. Therefore, elaboration occurs during run time. For example, the variable declarations that appear at the beginning of a Java method are elaborated when the method is called and the variables defined by those declarations are deallocated when the method completes its execution.

As their name indicates, stack-dynamic variables are allocated from the run-time stack.

Some languages—for example, C++ and Java—allow variable declarations to occur anywhere a statement can appear. In some implementations of these languages, all of the stack-dynamic variables declared in a function or method (not including those declared in nested blocks) may be bound to storage at the beginning of execution of the function or method, even though the declarations of some of these variables do not appear at the beginning. In such cases, the

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4. In some implementations, static variables are addressed through a base register, making accesses to them as costly as for stack-allocated variables.
variable becomes visible at the declaration, but the storage binding (and initialization, if it is specified in the declaration) occurs when the function or method begins execution. The fact that storage binding of a variable takes place before it becomes visible does not affect the semantics of the language.

The advantages of stack-dynamic variables are as follows: To be useful, at least in most cases, recursive subprograms require some form of dynamic local storage so that each active copy of the recursive subprogram has its own version of the local variables. These needs are conveniently met by stack-dynamic variables. Even in the absence of recursion, having stack-dynamic local storage for subprograms is not without merit, because all subprograms share the same memory space for their locals.

The disadvantages, relative to static variables, of stack-dynamic variables are the run-time overhead of allocation and deallocation, possibly slower accesses because indirect addressing is required, and the fact that subprograms cannot be history sensitive. The time required to allocate and deallocate stack-dynamic variables is not significant, because all of the stack-dynamic variables that are declared at the beginning of a subprogram are allocated and deallocated together, rather than by separate operations.

In Java, C++, and C#, variables defined in methods are by default stack dynamic.

All attributes other than storage are statically bound to stack-dynamic scalar variables. That is not the case for some structured types, as is discussed in Chapter 6. Implementation of allocation/deallocation processes for stack-dynamic variables is discussed in Chapter 10.

### 5.4.3.3 Explicit Heap-Dynamic Variables

**Explicit heap-dynamic variables** are nameless (abstract) memory cells that are allocated and deallocated by explicit run-time instructions written by the programmer. These variables, which are allocated from and deallocated to the heap, can only be referenced through pointer or reference variables. The heap is a collection of storage cells whose organization is highly disorganized due to the unpredictability of its use. The pointer or reference variable that is used to access an explicit heap-dynamic variable is created as any other scalar variable. An explicit heap-dynamic variable is created by either an operator (for example, in C++) or a call to a system subprogram provided for that purpose (for example, in C).

In C++, the allocation operator, named `new`, uses a type name as its operand. When executed, an explicit heap-dynamic variable of the operand type is created and its address is returned. Because an explicit heap-dynamic variable is bound to a type at compile time, that binding is static. However, such variables are bound to storage at the time they are created, which is during run time.

In addition to a subprogram or operator for creating explicit heap-dynamic variables, some languages include a subprogram or operator for explicitly destroying them.
As an example of explicit heap-dynamic variables, consider the following C++ code segment:

```cpp
int *intnode;  // Create a pointer
intnode = new int;  // Create the heap-dynamic variable

delete intnode;  // Deallocate the heap-dynamic variable
                 // to which intnode points
```

In this example, an explicit heap-dynamic variable of int type is created by the `new` operator. This variable can then be referenced through the pointer, `intnode`. Later, the variable is deallocated by the `delete` operator. C++ requires the explicit deallocation operator `delete`, because it does not use implicit storage reclamation, such as garbage collection.

In Java, all data except the primitive scalars are objects. Java objects are explicitly heap dynamic and are accessed through reference variables. Java has no way of explicitly destroying a heap-dynamic variable; rather, implicit garbage collection is used. Garbage collection is discussed in Chapter 6.

C# has both explicit heap-dynamic and stack-dynamic objects, all of which are implicitly deallocated. In addition, C# supports C++-style pointers. Such pointers are used to reference heap, stack, and even static variables and objects. These pointers have the same dangers as those of C++, and the objects they reference on the heap are not implicitly deallocated. Pointers are included in C# to allow C# components to interoperate with C and C++ components. To discourage their use, and also to make clear to any program reader that the code uses pointers, the header of any method that defines a pointer must include the reserved word `unsafe`.

Explicit heap-dynamic variables are often used to construct dynamic structures, such as linked lists and trees, that need to grow and/or shrink during execution. Such structures can be built conveniently using pointers or references and explicit heap-dynamic variables.

The disadvantages of explicit heap-dynamic variables are the difficulty of using pointer and reference variables correctly, the cost of references to the variables, and the complexity of the required storage management implementation. This is essentially the problem of heap management, which is costly and complicated. Implementation methods for explicit heap-dynamic variables are discussed at length in Chapter 6.

### 5.4.3.4 Implicit Heap-Dynamic Variables

Implicit heap-dynamic variables are bound to heap storage only when they are assigned values. In fact, all their attributes are bound every time they are assigned. For example, consider the following JavaScript assignment statement:

```javascript
highs = [74, 84, 86, 90, 71];
```

Regardless of whether the variable named `highs` was previously used in the program or what it was used for, it is now an array of five numeric values.
The advantage of such variables is that they have the highest degree of flexibility, allowing highly generic code to be written. One disadvantage of implicit heap-dynamic variables is the run-time overhead of maintaining all the dynamic attributes, which could include array subscript types and ranges, among others. Another disadvantage is the loss of some error detection by the compiler, as discussed in Section 5.4.2.2.

5.5 Scope

One of the important factors in understanding variables is scope. The scope of a variable is the range of statements in which the variable is visible. A variable is visible in a statement if it can be referenced or assigned in that statement.

The scope rules of a language determine how a particular occurrence of a name is associated with a variable, or in the case of a functional language, how a name is associated with an expression. In particular, scope rules determine how references to variables declared outside the currently executing subprogram or block are associated with their declarations and thus their attributes (blocks are discussed in Section 5.5.2). A clear understanding of these rules for a language is therefore essential to the ability to write or read programs in that language.

A variable is local in a program unit or block if it is declared there. The nonlocal variables of a program unit or block are those that are visible within the program unit or block but are not declared there. Global variables are a special category of nonlocal variables, which are discussed in Section 5.5.4.

Scoping issues of classes, packages, and namespaces are discussed in Chapter 11.

5.5.1 Static Scope

ALGOL 60 introduced the method of binding names to nonlocal variables called static scoping,⁵ which has been copied by many subsequent imperative languages and many nonimperative languages as well. Static scoping is so named because the scope of a variable can be statically determined—that is, prior to execution. This permits a human program reader (and a compiler) to determine the type of every variable in the program simply by examining its source code.

There are two categories of static-scoped languages: those in which subprograms can be nested, which creates nested static scopes, and those in which subprograms cannot be nested. In the latter category, static scopes are also created by subprograms but nested scopes are created only by nested class definitions and blocks.

Ada, JavaScript, Common Lisp, Scheme, Fortran 2003+, F#, and Python allow nested subprograms, but the C-based languages do not.

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⁵ Static scoping is sometimes called lexical scoping.
Our discussion of static scoping in this section focuses on those languages that allow nested subprograms. Initially, we assume that all scopes are associated with program units and that all referenced nonlocal variables are declared in other program units. In this chapter, it is assumed that scoping is the only method of accessing nonlocal variables in the languages under discussion. This is not true for all languages. It is not even true for all languages that use static scoping, but the assumption simplifies the discussion here.

When the reader of a program finds a reference to a variable, the attributes of the variable can be determined by finding the statement in which it is declared (either explicitly or implicitly). In static-scoped languages with nested subprograms, this process can be thought of in the following way. Suppose a reference is made to a variable \( x \) in subprogram \( \text{sub1} \). The correct declaration is found by first searching the declarations of subprogram \( \text{sub1} \). If no declaration is found for the variable there, the search continues in the declarations of the subprogram that declared subprogram \( \text{sub1} \), which is called its static parent. If a declaration of \( x \) is not found there, the search continues to the next-larger enclosing unit (the unit that declared \( \text{sub1} \)'s parent), and so forth, until a declaration for \( x \) is found or the largest unit’s declarations have been searched without success. In that case, an undeclared variable error is reported. The static parent of subprogram \( \text{sub1} \), and its static parent, and so forth up to and including the largest enclosing subprogram, are called the static ancestors of \( \text{sub1} \). Actual implementation techniques for static scoping, which are discussed in Chapter 10, are usually much more efficient than the process just described.

Consider the following JavaScript function, \( \text{big} \), in which the two functions \( \text{sub1} \) and \( \text{sub2} \) are nested:

```javascript
function big() {
  function sub1() {
    var x = 7;
    sub2();
  }
  function sub2() {
    var y = x;
  }
  var x = 3;
  sub1();
}
```

Under static scoping, the reference to the variable \( x \) in \( \text{sub2} \) is to the \( x \) declared in the procedure \( \text{big} \). This is true because the search for \( x \) begins in the procedure in which the reference occurs, \( \text{sub2} \), but no declaration for \( x \) is found there. The search continues in the static parent of \( \text{sub2}, \text{big} \), where the declaration of \( x \) is found. The \( x \) declared in \( \text{sub1} \) is ignored, because it is not in the static ancestry of \( \text{sub2} \).

---

6. Nonlocal variables not defined in other program units are discussed in Section 5.5.4.
In some languages that use static scoping, regardless of whether nested subprograms are allowed, some variable declarations can be hidden from some other code segments. For example, consider again the JavaScript function `big`. The variable `x` is declared in both `big` and in `sub1`, which is nested inside `big`. Within `sub1`, every simple reference to `x` is to the local `x`. Therefore, the outer `x` is hidden from `sub1`.

### 5.5.2 Blocks

Many languages allow new static scopes to be defined in the midst of executable code. This powerful concept, introduced in ALGOL 60, allows a section of code to have its own local variables whose scope is minimized. Such variables are typically stack dynamic, so their storage is allocated when the section is entered and deallocated when the section is exited. Such a section of code is called a block. Blocks provide the origin of the phrase **block-structured language**.

The C-based languages allow any compound statement (a statement sequence surrounded by matched braces) to have declarations and thereby define a new scope. Such compound statements are called blocks. For example, if `list` were an integer array, one could write the following:

```c
if (list[i] < list[j]) {
    int temp;
    temp = list[i];
    list[i] = list[j];
    list[j] = temp;
}
```

The scopes created by blocks, which could be nested in larger blocks, are treated exactly like those created by subprograms. References to variables in a block that are not declared there are connected to declarations by searching enclosing scopes (blocks or subprograms) in order of increasing size.

Consider the following skeletal C function:

```c
void sub() {
    int count;
    ...
    while (. . .) {
        int count;
        count++;
        ...
    }
    ...
}
```

The reference to `count` in the `while` loop is to that loop’s local `count`. In this case, the `count` of `sub` is hidden from the code inside the `while` loop. In general, a declaration for a variable effectively hides any declaration of a variable with
the same name in a larger enclosing scope.\footnote{As discussed in Section 5.5.4, in C++, such hidden global variables can be accessed in the inner scope using the scope operator (::).} Note that this code is legal in C and C++ but illegal in Java and C#. The designers of Java and C# believed that the reuse of names in nested blocks was too error prone to be allowed.

Although JavaScript uses static scoping for its nested functions, nonfunction blocks cannot be defined in the language.

Most functional programming languages include a construct that is related to the blocks of the imperative languages, usually named \textit{let}. These constructs have two parts, the first of which is to bind names to values, usually specified as expressions. The second part is an expression that uses the names defined in the first part. Programs in functional languages are comprised of expressions, rather than statements. Therefore, the final part of a \textit{let} construct is an expression, rather than a statement. In Scheme, a \textit{let} construct is a call to the function \texttt{LET} with the following form:

\begin{verbatim}
(LET ( 
    (name\textsubscript{1} expression\textsubscript{1})
    . . .
    (name\textsubscript{n} expression\textsubscript{n}))
  expression)
\end{verbatim}

The semantics of the call to \texttt{LET} is as follows: The first \textit{n} expressions are evaluated and the values are assigned to the associated names. Then, the final expression is evaluated and the return value of \texttt{LET} is that value. This differs from a block in an imperative language in that the names are of values; they are not variables in the imperative sense. Once set, they cannot be changed. However, they are like local variables in a block in an imperative language in that their scope is local to the call to \texttt{LET}. Consider the following call to \texttt{LET}:

\begin{verbatim}
(LET ( 
    (top (+ a b))
    (bottom (- c d)))
  (/ top bottom))
\end{verbatim}

This call computes and returns the value of the expression \((a + b) / (c - d)\).

In ML, the form of a \textit{let} construct is as follows:

\begin{verbatim}
let 
  val name\textsubscript{1} = expression\textsubscript{1} 
  . . .
  val name\textsubscript{n} = expression\textsubscript{n} 
in 
  expression 
end;
\end{verbatim}
5.5 Scope

Each `val` statement binds a name to an expression. As with Scheme, the names in the first part are like the named constants of imperative languages; once set, they cannot be changed.\(^8\) Consider the following `let` construct:

```fsharp
let
    val top = a + b
    val bottom = c - d
in
    top / bottom
end;
```

The general form of a `let` construct in F# is as follows:

```fsharp
let left_side = expression
```

The left_side of `let` can be a name or a tuple pattern (a sequence of names separated by commas).

The scope of a name defined with `let` inside a function definition is from the end of the defining expression to the end of the function. The scope of `let` can be limited by indenting the following code, which creates a new local scope. Although any indentation will work, the convention is that the indentation is four spaces. Consider the following code:

```fsharp
let n1 =
    let n2 = 7
    let n3 = n2 + 3
    n3;;
let n4 = n3 + n1;;
```

The scope of `n1` extends over all of the code. However, the scope of `n2` and `n3` ends when the indentation ends. So, the use of `n3` in the last `let` causes an error. The last line of the `let n1` scope is the value bound to `n1`; it could be any expression.

Chapter 15 includes more details of the let constructs in Scheme, ML, Haskell, and F#.

5.5.3 Declaration Order

In C89, as well as in some other languages, all data declarations in a function except those in nested blocks must appear at the beginning of the function. However, some languages—for example, C99, C++, Java, JavaScript, and C#—allow variable declarations to appear anywhere a statement can appear in a program unit. Declarations may create scopes that are not associated with compound statements or subprograms. For example, in C99, C++, and Java, the scope of all local variables is from their declarations to the ends of the blocks in which those declarations appear.

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\(^8\) In Chapter 15, we will see that they can be reset, but that the process actually creates a new name.
In C#, the scope of any variable declared in a block is the whole block, regardless of the position of the declaration in the block, as long as it is not in a nested block. The same is true for methods. Recall that C# does not allow the declaration of a variable in a nested block to have the same name as a variable in a nesting scope. For example, consider the following C# code:

```csharp
{ int x;
  ...
  { int x; // Illegal
    ...
  }
  ...
}
```

Because the scope of a declaration is the whole block, the following nested declaration of x is also illegal:

```csharp
{
  { int x; // Illegal
    ...
  }
  int x;
}
```

Note that C# still requires that all be declared before they are used. Therefore, although the scope of a variable extends from the declaration to the top of the block or subprogram in which that declaration appears, the variable still cannot be used above its declaration.

In JavaScript, local variables can be declared anywhere in a function, but the scope of such a variable is always the entire function. If used before its declaration in the function, such a variable has the value `undefined`. The reference is not illegal.

The for statements of C++, Java, and C# allow variable definitions in their initialization expressions. In early versions of C++, the scope of such a variable was from its definition to the end of the smallest enclosing block. In the standard version, however, the scope is restricted to the for construct, as is the case with Java and C#. Consider the following skeletal method:

```csharp
void fun() {
  ...
  for (int count = 0; count < 10; count++){
    ...
  }
  ...
}
```
In later versions of C++, as well as in Java and C#, the scope of count is from the for statement to the end of its body (the right brace).

5.5.4 Global Scope

Some languages, including C, C++, PHP, JavaScript, and Python, allow a program structure that is a sequence of function definitions, in which variable definitions can appear outside the functions. Definitions outside functions in a file create global variables, which potentially can be visible to those functions.

C and C++ have both declarations and definitions of global data. Declarations specify types and other attributes but do not cause allocation of storage. Definitions specify attributes and cause storage allocation. For a specific global name, a C program can have any number of compatible declarations, but only a single definition.

A declaration of a variable outside function definitions specifies that the variable is defined in a different file. A global variable in C is implicitly visible in all subsequent functions in the file, except those that include a declaration of a local variable with the same name. A global variable that is defined after a function can be made visible in the function by declaring it to be external, as in the following:

```
extern int sum;
```

In C99, definitions of global variables usually have initial values. Declarations of global variables never have initial values. If the declaration is outside function definitions, it need not include the extern qualifier.

This idea of declarations and definitions carries over to the functions of C and C++, where prototypes declare names and interfaces of functions but do not provide their code. Function definitions, on the other hand, are complete.

In C++, a global variable that is hidden by a local with the same name can be accessed using the scope operator (::). For example, if x is a global that is hidden in a function by a local named x, the global could be referenced as ::x.

PHP statements can be interspersed with function definitions. Variables in PHP are implicitly declared when they appear in statements. Any variable that is implicitly declared outside any function is a global variable; variables implicitly declared in functions are local variables. The scope of global variables extends from their declarations to the end of the program but skips over any subsequent function definitions. So, global variables are not implicitly visible in any function. Global variables can be made visible in functions in their scope in two ways: (1) If the function includes a local variable with the same name as a global, that global can be accessed through the $GLOBALS array, using the name of the global as a string literal subscript, and (2) if there is no local variable in the function with the same name as the global, the global can be
made visible by including it in a global declaration statement. Consider the following example:

```php
$day = "Monday";
$month = "January";

function calendar() {
  $day = "Tuesday";
  global $month;
  print "local day is $day 
";
  $gday = $GLOBALS['day'];
  print "global day is $gday <br ">
  print "global month is $month 
";
}
calendar();
```

Interpretation of this code produces the following:

```
local day is Tuesday
global day is Monday
global month is January
```

The global variables of JavaScript are very similar to those of PHP, except that there is no way to access a global variable in a function that has declared a local variable with the same name.

The visibility rules for global variables in Python are unusual. Variables are not normally declared, as in PHP. They are implicitly declared when they appear as the targets of assignment statements. A global variable can be referenced in a function, but a global variable can be assigned in a function only if it has been declared to be global in the function. Consider the following examples:

```python
day = "Monday"

def tester():
  print "The global day is:", day

tester()
```

The output of this script, because globals can be referenced directly in functions, is as follows:

```
The global day is: Monday
```

The following script attempts to assign a new value to the global day:
5.5 Scope

```python
day = "Monday"

def tester():
    print "The global day is:", day
day = "Tuesday"
    print "The new value of day is:", day
tester()
```

This script creates an `UnboundLocalError` error message, because the assignment to `day` in the second line of the body of the function makes `day` a local variable, which makes the reference to `day` in the first line of the body of the function an illegal forward reference to the local.

The assignment to `day` can be to the global variable if `day` is declared to be global at the beginning of the function. This prevents the assignment to `day` from creating a local variable. This is shown in the following script:

```python
day = "Monday"

def tester():
    global day
    print "The global day is:", day
day = "Tuesday"
    print "The new value of day is:", day
tester()
```

The output of this script is as follows:

```
The global day is: Monday
The new value of day is: Tuesday
```

Functions can be nested in Python. Variables defined in nesting functions are accessible in a nested function through static scoping, but such variables must be declared `nonlocal` in the nested function. An example skeletal program in Section 5.7 illustrates accesses to nonlocal variables.

All names defined outside function definitions in F# are globals. Their scope extends from their definitions to the end of the file.

Declaration order and global variables are also issues in the class and member declarations in object-oriented languages. These are discussed in Chapter 12.

---

9. The `nonlocal` reserved word was introduced in Python 3.
5.5.5 Evaluation of Static Scoping

Static scoping provides a method of nonlocal access that works well in many situations. However, it is not without its problems. First, in most cases it allows more access to both variables and subprograms than is necessary. It is simply too crude a tool for concisely specifying such restrictions. Second, and perhaps more important, is a problem related to program evolution. Software is highly dynamic—programs that are used regularly continually change. These changes often result in restructuring, thereby destroying the initial structure that restricted variable and subprogram access in a static-scoped language. To avoid the complexity of maintaining these access restrictions, developers often discard structure when it gets in the way. Thus, getting around the restrictions of static scoping can lead to program designs that bear little resemblance to the original, even in areas of the program in which changes have not been made. Designers are encouraged to use far more globals than are necessary. All subprograms can end up being nested at the same level, in the main program, using globals instead of deeper levels of nesting. Moreover, the final design may be awkward and contrived, and it may not reflect the underlying conceptual design. These and other defects of static scoping are discussed in detail in Clarke, Wileden, and Wolf (1980). An alternative to the use of static scoping to control access to variables and subprograms is an encapsulation construct, which is included in many newer languages. Encapsulation constructs are discussed in Chapter 11.

5.5.6 Dynamic Scope

The scope of variables in APL, SNOBOL4, and the early versions of Lisp is dynamic. Perl and Common Lisp also allow variables to be declared to have dynamic scope, although the default scoping mechanism in these languages is static. Dynamic scoping is based on the calling sequence of subprograms, not on their spatial relationship to each other. Thus, the scope can be determined only at run time.

Consider again the function `big` from Section 5.5.1, which is reproduced here, minus the function calls:

```plaintext
function big() {
    function sub1() {
        var x = 7;
    }
    function sub2() {
        var y = x;
        var z = 3;
    }
    var x = 3;
}
```

10. Sounds like the structure of a C program, doesn’t it?
Assume that dynamic-scoping rules apply to nonlocal references. The meaning of the identifier \( x \) referenced in \texttt{sub2} is dynamic—it cannot be determined at compile time. It may reference the variable from either declaration of \( x \), depending on the calling sequence.

One way the correct meaning of \( x \) can be determined during execution is to begin the search with the local declarations. This is also the way the process begins with static scoping, but that is where the similarity between the two techniques ends. When the search of local declarations fails, the declarations of the dynamic parent, or calling function, are searched. If a declaration for \( x \) is not found there, the search continues in that function’s dynamic parent, and so forth, until a declaration for \( x \) is found. If none is found in any dynamic ancestor, it is a run-time error.

Consider the two different call sequences for \texttt{sub2} in the earlier example. First, \texttt{big} calls \texttt{sub1}, which calls \texttt{sub2}. In this case, the search proceeds from the local procedure, \texttt{sub2}, to its caller, \texttt{sub1}, where a declaration for \( x \) is found. So, the reference to \( x \) in \texttt{sub2} in this case is to the \( x \) declared in \texttt{sub1}. Next, \texttt{sub2} is called directly from \texttt{big}. In this case, the dynamic parent of \texttt{sub2} is \texttt{big}, and the reference is to the \( x \) declared in \texttt{big}.

Note that if static scoping were used, in either calling sequence discussed, the reference to \( x \) in \texttt{sub2} would be to \texttt{big}'s \( x \).

Perl’s dynamic scoping is unusual—in fact, it is not exactly like that discussed in this section, although the semantics are often that of traditional dynamic scoping (see Programming Exercise 1).

### 5.5.7 Evaluation of Dynamic Scoping

The effect of dynamic scoping on programming is profound. When dynamic scoping is used, the correct attributes of nonlocal variables visible to a program statement cannot be determined statically. Furthermore, a reference to the name of such a variable is not always to the same variable. A statement in a subprogram that contains a reference to a nonlocal variable can refer to different nonlocal variables during different executions of the subprogram. Several kinds of programming problems follow directly from dynamic scoping.

First, during the time span beginning when a subprogram begins its execution and ending when that execution ends, the local variables of the subprogram are all visible to any other executing subprogram, regardless of its textual proximity or how execution got to the currently executing subprogram. There is no way to protect local variables from this accessibility. Subprograms are always executed in the environment of all previously called subprograms that have not yet completed their executions. As a result, dynamic scoping results in less reliable programs than static scoping.

A second problem with dynamic scoping is the inability to type check references to nonlocals statically. This problem results from the inability to statically find the declaration for a variable referenced as a nonlocal.

Dynamic scoping also makes programs much more difficult to read, because the calling sequence of subprograms must be known to determine the
meaning of references to nonlocal variables. This task can be virtually impossible for a human reader.

Finally, accesses to nonlocal variables in dynamic-scoped languages take far longer than accesses to nonlocals when static scoping is used. The reason for this is explained in Chapter 10.

On the other hand, dynamic scoping is not without merit. In many cases, the parameters passed from one subprogram to another are variables that are defined in the caller. None of these needs to be passed in a dynamically scoped language, because they are implicitly visible in the called subprogram.

It is not difficult to understand why dynamic scoping is not as widely used as static scoping. Programs in static-scoped languages are easier to read, more reliable, and execute faster than equivalent programs in dynamic-scoped languages. It was precisely for these reasons that dynamic scoping was replaced by static scoping in most current dialects of Lisp. Implementation methods for both static and dynamic scoping are discussed in Chapter 10.

5.6 Scope and Lifetime

Sometimes the scope and lifetime of a variable appear to be related. For example, consider a variable that is declared in a Java method that contains no method calls. The scope of such a variable is from its declaration to the end of the method. The lifetime of that variable is the period of time beginning when the method is entered and ending when execution of the method terminates. Although the scope and lifetime of the variable appear to be related in this case.

This apparent relationship between scope and lifetime does not hold in other situations. In C and C++, for example, a variable that is declared in a function using the specifier `static` is statically bound to the scope of that function and is also statically bound to storage. So, its scope is static and local to the function, but its lifetime extends over the entire execution of the program of which it is a part.

Scope and lifetime are also unrelated when subprogram calls are involved. Consider the following C++ functions:

```cpp
void printhead() {
    . . .
} /* end of printhead */
void compute() {
    int sum;
    . . .
    printhead();
} /* end of compute */
```

The scope of the variable `sum` is completely contained within the `compute` function. It does not extend to the body of the function `printhead`, although `printhead` executes in the midst of the execution of `compute`. However, the
lifetime of \texttt{sum} extends over the time during which \texttt{printhead}er executes. Whatever storage location \texttt{sum} is bound to before the call to \texttt{printhead}er, that binding will continue during and after the execution of \texttt{printhead}er.

### 5.7 Referencing Environments

The \textbf{referencing environment} of a statement is the collection of all variables that are visible in the statement. The referencing environment of a statement in a static-scoped language is the variables declared in its local scope plus the collection of all variables of its ancestor scopes that are visible. In such a language, the referencing environment of a statement is needed while that statement is being compiled, so code and data structures can be created to allow references to variables from other scopes during run time. Techniques for implementing references to nonlocal variables in both static- and dynamic-scoped languages are discussed in Chapter 10.

In Python, scopes can be created by function definitions. The referencing environment of a statement includes the local variables, plus all of the variables declared in the functions in which the statement is nested (excluding variables in nonlocal scopes that are hidden by declarations in nearer functions). Each function definition creates a new scope and thus a new environment. Consider the following Python skeletal program:

```python
g = 3; # A global

def sub1():
a = 5; # Creates a local
b = 7; # Creates another local
. . . <--------------------------------------------- 1

def sub2():
global g; # Global g is now assignable here
c = 9; # Creates a new local
. . . <--------------------------------------------- 2

def sub3():
    nonlocal c: # Makes nonlocal c visible here
g = 11; # Creates a new local
. . . <--------------------------------------------- 3
```

The referencing environments of the indicated program points are as follows:

<table>
<thead>
<tr>
<th>Point</th>
<th>Referencing Environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>local a and b (of sub1), global g for reference, but not for assignment</td>
</tr>
<tr>
<td>2</td>
<td>local c (of sub2), global g for both reference and for assignment</td>
</tr>
<tr>
<td>3</td>
<td>nonlocal c (of sub2), local g (of sub3)</td>
</tr>
</tbody>
</table>
Now consider the variable declarations of this skeletal program. First, note that, although the scope of \texttt{sub1} is at a higher level (it is less deeply nested) than \texttt{sub3}, the scope of \texttt{sub1} is not a static ancestor of \texttt{sub3}, so \texttt{sub3} does not have access to the variables declared in \texttt{sub1}. There is a good reason for this. The variables declared in \texttt{sub1} are stack dynamic, so they are not bound to storage if \texttt{sub1} is not in execution. Because \texttt{sub3} can be in execution when \texttt{sub1} is not, it cannot be allowed to access variables in \texttt{sub1}, which would not necessarily be bound to storage during the execution of \texttt{sub3}.

A subprogram is \textbf{active} if its execution has begun but has not yet terminated. The referencing environment of a statement in a dynamically scoped language is the locally declared variables, plus the variables of all other subprograms that are currently active. Once again, some variables in active subprograms can be hidden from the referencing environment. Recent subprogram activations can have declarations for variables that hide variables with the same names in previous subprogram activations.

Consider the following example program. Assume that the only function calls are the following: \texttt{main} calls \texttt{sub2}, which calls \texttt{sub1}.

\begin{verbatim}
void sub1() {
    int a, b;
    . . . <--------- 1
} /* end of sub1 */

void sub2() {
    int b, c;
    . . . <--------- 2
    sub1();
} /* end of sub2 */

void main() {
    int c, d;
    . . . <--------- 3
    sub2();
} /* end of main */
\end{verbatim}

The referencing environments of the indicated program points are as follows:

\begin{center}
\begin{tabular}{ll}
\textbf{Point} & \textbf{Referencing Environment} \\
1 & \texttt{a} and \texttt{b} of \texttt{sub1}, \texttt{c} of \texttt{sub2}, \texttt{d} of \texttt{main}, (\texttt{c} of \texttt{main} and \texttt{b} of \texttt{sub2} are hidden) \\
2 & \texttt{b} and \texttt{c} of \texttt{sub2}, \texttt{d} of \texttt{main}, (\texttt{c} of \texttt{main} is hidden) \\
3 & \texttt{c} and \texttt{d} of \texttt{main} \\
\end{tabular}
\end{center}

\section{5.8 Named Constants}

A \textbf{named constant} is a variable that is bound to a value only once. Named constants are useful as aids to readability and program reliability. Readability can be improved, for example, by using the name \texttt{pi} instead of the constant 3.14159265.
Another important use of named constants is to parameterize a program. For example, consider a program that processes a fixed number of data values, say 100. Such a program usually uses the constant 100 in a number of locations for declaring array subscript ranges and for loop control limits. Consider the following skeletal Java program segment:

```java
void example() {
    int[] intList = new int[100];
    String[] strList = new String[100];
    . . .
    for (index = 0; index < 100; index++) {
        . . .
    }
    . . .
    for (index = 0; index < 100; index++) {
        . . .
    }
    . . .
    average = sum / 100;
    . . .
}
```

When this program must be modified to deal with a different number of data values, all occurrences of 100 must be found and changed. On a large program, this can be tedious and error prone. An easier and more reliable method is to use a named constant as a program parameter, as follows:

```java
void example() {
    final int len = 100;
    int[] intList = new int[len];
    String[] strList = new String[len];
    . . .
    for (index = 0; index < len; index++) {
        . . .
    }
    . . .
    for (index = 0; index < len; index++) {
        . . .
    }
    . . .
    average = sum / len;
    . . .
}
```

Now, when the length must be changed, only one line must be changed (the variable len), regardless of the number of times it is used in the program. This is another example of the benefits of abstraction. The name len is an abstraction for the number of elements in some arrays and the number of iterations in some loops. This illustrates how named constants can aid modifiability.
C++ allows dynamic binding of values to named constants. This allows expressions containing variables to be assigned to constants in the declarations. For example, the C++ statement

```cpp
const int result = 2 * width + 1;
```

declares result to be an integer type named constant whose value is set to the value of the expression $2 \times width + 1$, where the value of the variable width must be visible when result is allocated and bound to its value.

Java also allows dynamic binding of values to named constants. In Java, named constants are defined with the `final` reserved word (as in the earlier example). The initial value can be given in the declaration statement or in a subsequent assignment statement. The assigned value can be specified with any expression.

C# has two kinds of named constants: those defined with `const` and those defined with `readonly`. The `const` named constants, which are implicitly `static`, are statically bound to values; that is, they are bound to values at compile time, which means those values can be specified only with literals or other `const` members. The `readonly` named constants, which are dynamically bound to values, can be assigned in the declaration or with a static constructor.11 So, if a program needs a constant-valued object whose value is the same on every use of the program, a `const` constant is used. However, if a program needs a constant-valued object whose value is determined only when the object is created and can be different for different executions of the program, then a `readonly` constant is used.

The discussion of binding values to named constants naturally leads to the topic of initialization, because binding a value to a named constant is the same process, except it is permanent.

In many instances, it is convenient for variables to have values before the code of the program or subprogram in which they are declared begins executing. The binding of a variable to a value at the time it is bound to storage is called `initialization`. If the variable is statically bound to storage, binding and initialization occur before run time. In these cases, the initial value must be specified as a literal or an expression whose only nonliteral operands are named constants that have already been defined. If the storage binding is dynamic, initialization is also dynamic and the initial values can be any expression.

In most languages, initialization is specified on the declaration that creates the variable. For example, in C++, we could have

```cpp
int sum = 0;
int* ptrSum = &sum;
char name[] = "George Washington Carver";
```

---

11. Static constructors in C# run at some indeterminate time before the class is instantiated.
SUMMARY

Case sensitivity and the use of underscores are the design issues for names. Variables can be characterized by the sextuple of attributes: name, address, value, type, lifetime, and scope.

Aliases are two or more variables bound to the same storage address. They are regarded as detrimental to reliability but are difficult to eliminate entirely from a language.

Binding is the association of attributes with program entities. Knowledge of the binding times of attributes to entities is essential to understanding the semantics of programming languages. Binding can be static or dynamic. Declarations, either explicit or implicit, provide a means of specifying the static binding of variables to types. In general, dynamic binding allows greater flexibility but at the expense of readability, efficiency, and reliability.

Scalar variables can be separated into four categories by considering their lifetimes: static, stack dynamic, explicit heap dynamic, and implicit heap dynamic.

Static scoping is a central feature of ALGOL 60 and some of its descendants. It provides a simple, reliable, and efficient method of allowing visibility of non-local variables in subprograms. Dynamic scoping provides more flexibility than static scoping but, again, at the expense of readability, reliability, and efficiency.

Most functional languages allow the user to create local scopes with let constructs, which limit the scope of their defined names.

The referencing environment of a statement is the collection of all of the variables that are visible to that statement.

Named constants are simply variables that are bound to values only once.

REVIEW QUESTIONS

1. What is a reserved word?
2. In Java and C#, how long can a name be?
3. What is the address of a variable?
4. How can a variable be characterized?
5. What is type inference? Give an example.
6. What is deallocation of a memory cell?
7. After language design and implementation, what are the four times bindings can take place in a program?
8. What is the lifetime of a variable?
9. What is the use of a stack-dynamic variable?
10. What are the advantages and disadvantages of dynamic type binding?
11. Define static, stack-dynamic, explicit heap-dynamic, and implicit heap-dynamic variables. What are their advantages and disadvantages?
12. What is a block-structured language?
13. How is a reference to a nonlocal variable in a static-scoped program connected to its definition?
14. What is the general problem with static scoping?
15. What is the referencing environment of a statement?
16. What is a static ancestor of a subprogram? What is a dynamic ancestor of a subprogram?
17. What is a block?
18. What is the purpose of the \texttt{let} constructs in functional languages?
19. What is the difference between the names defined in an ML \texttt{let} construct from the variables declared in a C block?
20. Describe the encapsulation of an F# \texttt{let} inside a function and outside all functions.
21. When can we call a subprogram active?
22. What is variable initialization?

\section*{Problem Set}

1. Which of the following identifier forms is most readable? Support your decision.
   \begin{itemize}
   \item \texttt{FirstName}
   \item \texttt{First_Name}
   \item \texttt{firstname}
   \end{itemize}
2. Some programming languages have static variables. What are the obvious advantages and disadvantages of having static variables?
3. Write a simple assignment statement with one arithmetic operator in some language you know. For each component of the statement, list the various bindings that are required to determine the semantics when the statement is executed. For each binding, indicate the binding time used for the language.
4. Dynamic type binding is closely related to implicit heap-dynamic variables. Explain this relationship.
5. Describe a situation when dynamic scoping makes a program difficult to read.
6. Consider the following JavaScript skeletal program:
   \begin{verbatim}
   // The main program
   var x;
   function sub1() {
   \end{verbatim}
\begin{verbatim}
var x;
function sub2() {
    ...
}
}
function sub3() {
    ...
}

Assume that the execution of this program is in the following unit order:
main calls sub1
sub1 calls sub2
sub2 calls sub3

a. Assuming static scoping, in the following, which declaration of x is
   the correct one for a reference to x?
   i. sub1
   ii. sub2
   iii. sub3

b. Repeat part a, but assume dynamic scoping.

7. Assume the following JavaScript program was interpreted using
   static-scoping rules. What value of x is displayed in function sub1?
   Under dynamic-scoping rules, what value of x is displayed in function
   sub1?

\begin{verbatim}
var x;
function sub1() {
    document.write("x = " + x + ");
}

function sub2() {
    var x;
    x = 10;
    sub1();
}

x = 5;
sub2();
\end{verbatim}

8. Consider the following JavaScript program:

\begin{verbatim}
var x, y, z;
function sub1() {
    var a, y, z;
\end{verbatim}
function sub2() {
    var a, b, z;
    ...
}
...
}

function sub3() {
    var a, x, w;
    ...
}

List all the variables, along with the program units where they are declared, that are visible in the bodies of sub1, sub2, and sub3, assuming static scoping is used.

9. Consider the following Python program:

```python
x = 1;
y = 3;
z = 5;
def sub1():
a = 7;
y = 9;
z = 11;
...
def sub2():
global x;
a = 13;
x = 15;
w = 17;
...
def sub3():
    nonlocal a;
a = 19;
b = 21;
z = 23;
...
```

List all the variables, along with the program units where they are declared, that are visible in the bodies of sub1, sub2, and sub3, assuming static scoping is used.
10. Consider the following C program:

```c
void fun(void) {
    int a, b, c; /* definition 1 */
    . . .
    while (. . .) {
        int b, c, d; /* definition 2 */
        . . . <------------- 1
        while (. . .) {
            int c, d, e; /* definition 3 */
            . . . <------------- 2
        }
        . . . <------------- 3
    }
    . . . <------------- 4
}
```

For each of the four marked points in this function, list each visible variable, along with the number of the definition statement that defines it.

11. Consider the following skeletal C program:

```c
void fun1(void); /* prototype */
void fun2(void); /* prototype */
void fun3(void); /* prototype */
void main() {
    int a, b, c;
    . . .
}
void fun1(void) {
    int b, c, d;
    . . .
}
void fun2(void) {
    int c, d, e;
    . . .
}
void fun3(void) {
    int d, e, f;
    . . .
}
```
Given the following calling sequences and assuming that dynamic scoping is used, what variables are visible during execution of the last function called? Include with each visible variable the name of the function in which it was defined.

a. main calls fun1; fun1 calls fun2; fun2 calls fun3.
b. main calls fun1; fun1 calls fun3.
c. main calls fun2; fun2 calls fun3; fun3 calls fun1.
d. main calls fun3; fun3 calls fun1.
e. main calls fun1; fun1 calls fun3; fun3 calls fun2.
f. main calls fun3; fun3 calls fun2; fun2 calls fun1.

12. Consider the following program, written in JavaScript-like syntax:

```javascript
// main program
var x, y, z;

function sub1() {
    var a, y, z;
    ...
}
function sub2() {
    var a, b, z;
    ...
}
function sub3() {
    var a, x, w;
    ...
}
```

Given the following calling sequences and assuming that dynamic scoping is used, what variables are visible during execution of the last subprogram activated? Include with each visible variable the name of the unit where it is declared.

a. main calls sub1; sub1 calls sub2; sub2 calls sub3.
b. main calls sub1; sub1 calls sub3.
c. main calls sub2; sub2 calls sub3; sub3 calls sub1.
d. main calls sub3; sub3 calls sub1.
e. main calls sub1; sub1 calls sub3; sub3 calls sub2.
f. main calls sub3; sub3 calls sub2; sub2 calls sub1.
1. Perl allows both static and a kind of dynamic scoping. Write a Perl program that uses both and clearly shows the difference in effect of the two. Explain clearly the difference between the dynamic scoping described in this chapter and that implemented in Perl.

2. Write a Common Lisp program that clearly shows the difference between static and dynamic scoping.

3. Write a JavaScript script that has subprograms nested three deep and in which each nested subprogram references variables defined in all of its enclosing subprograms.

4. Repeat Programming Exercise 3 with Python.

5. Write a C99 function that includes the following sequence of statements:

   ```c
   x = 21;
   int x;
   x = 42;
   ```

   Run the program and explain the results. Rewrite the same code in C++ and Java and compare the results.

6. Write test programs in C++, Java, and C# to determine the scope of a variable declared in a `for` statement. Specifically, the code must determine whether such a variable is visible after the body of the `for` statement.

7. Write three functions in C or C++: one that declares a large array statically, one that declares the same large array on the stack, and one that creates the same large array from the heap. Call each of the subprograms a large number of times (at least 100,000) and output the time required by each. Explain the results.
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6.1 Introduction
6.2 Primitive Data Types
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6.12 Type Checking
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6.15 Theory and Data Types
This chapter first introduces the concept of a data type and the characteristics of the common primitive data types. Then, the designs of enumeration and subrange types are discussed. Next, the details of structured data types—specifically arrays, associative arrays, records, tuples, lists, and unions—are investigated. This section is followed by an in-depth look at pointers and references.

For each of the various categories of data types, the design issues are stated and the design choices made by the designers of some common languages are described. These designs are then evaluated.

The next three sections provide a thorough investigation of type checking, strong typing, and type equivalence rules. The last section of the chapter briefly introduces the basics of the theory of data types.

Implementation methods for data types sometimes have a significant impact on their design. Therefore, implementation of the various data types is another important part of this chapter, especially arrays.

6.1 Introduction

A data type defines a collection of data values and a set of predefined operations on those values. Computer programs produce results by manipulating data. An important factor in determining the ease with which they can perform this task is how well the data types available in the language being used match the objects in the real world of the problem being addressed. Therefore, it is crucial that a language supports an appropriate collection of data types and structures.

The contemporary concepts of data typing have evolved over the last 60 years. In the earliest languages, all problem space data structures had to be modeled with only a few basic language-supported data structures. For example, in pre-90 Fortrans, linked lists and binary trees were implemented with arrays.

The data structures of COBOL took the first step away from the Fortran I model by allowing programmers to specify the accuracy of decimal data values, and also by providing a structured data type for records of information. PL/I extended the capability of accuracy specification to integer and floating-point types. The designers of PL/I included many data types, with the intent of supporting a large range of applications. A better approach, introduced in ALGOL 68, is to provide a few basic types and a few flexible structure-defining operators that allow a programmer to design a data structure for each need. Clearly, this was one of the most important advances in the evolution of data type design. User-defined types also provide improved readability through the use of meaningful names for types. They allow type checking of the variables of a special category of use, which would otherwise not be possible. User-defined types also aid modifiability: A programmer can change the type of a category of variables in a program by changing a type definition statement only.

Taking the concept of a user-defined type a step further, we arrive at abstract data types, which are supported by most programming languages designed since the mid-1980s. The fundamental idea of an abstract data type
is that the interface of a type, which is visible to the user, is separated from the representation and set of operations on values of that type, which are hidden from the user. All of the types provided by a high-level programming language are abstract data types. User-defined abstract data types are discussed in detail in Chapter 11.

There are a number of uses of the type system of a programming language. The most practical of these is error detection. The process and value of type checking, which is directed by the type system of the language, are discussed in Section 6.12. A second use of a type system is the assistance it provides for program modularization. This results from the cross-module type checking that ensures the consistency of the interfaces among modules. Another use of a type system is documentation. The type declarations in a program document information about its data, which provides clues about the program’s behavior.

The type system of a programming language defines how a type is associated with each expression in the language and includes its rules for type equivalence and type compatibility. Certainly, one of the most important parts of understanding the semantics of a programming language is understanding its type system.

The two most common structured (nonscalar) data types in the imperative languages are arrays and records, although the popularity of associative arrays has increased significantly in recent years. Lists have been a central part of functional programming languages since the first such language appeared in 1959 (Lisp). Over the last decade, the increasing popularity of functional programming has led to lists being added to primarily imperative languages, such as Python and C#.

The structured data types are defined with type operators, or constructors, which are used to form type expressions. For example, C uses brackets and asterisks as type operators to specify arrays and pointers.

It is convenient, both logically and concretely, to think of variables in terms of descriptors. A descriptor is the collection of the attributes of a variable. In an implementation, a descriptor is an area of memory that stores the attributes of a variable. If the attributes are all static, descriptors are required only at compile time. These descriptors are built by the compiler, usually as a part of the symbol table, and are used during compilation. For dynamic attributes, however, part or all of the descriptor must be maintained during execution. In this case, the descriptor is used by the run-time system. In all cases, descriptors are used for type checking and building the code for the allocation and deallocation operations.

Care must be taken when using the term variable. One who uses only traditional imperative languages may think of identifiers as variables, but that can lead to confusion when considering data types. Identifiers do not have data types in some programming languages. It is wise to remember that identifiers are just one of the attributes of a variable.

The word object is often associated with the value of a variable and the space it occupies. In this book, however, we reserve object exclusively for instances of user-defined and language-defined abstract data types, rather than for the
values of all program variables of predefined types. Objects are discussed in
detail in Chapters 11 and 12.

In the following sections, many common data types are discussed. For most,
design issues particular to the type are stated. For all, one or more example
designs are described. One design issue is fundamental to all data types: What
operations are provided for variables of the type, and how are they specified?

6.2 Primitive Data Types

Data types that are not defined in terms of other types are called **primitive
data types**. Nearly all programming languages provide a set of primitive data
types. Some of the primitive types are merely reflections of the hardware—for
example, most integer types. Others require only a little nonhardware support
for their implementation.

To specify the structured types, the primitive data types of a language are
used, along with one or more type constructors.

6.2.1 Numeric Types

Some early programming languages only had numeric primitive types. Numeric
types still play a central role among the collections of types supported by con-
temporary languages.

6.2.1.1 Integer

The most common primitive numeric data type is **integer**. The hardware
of many computers supports several sizes of integers. These sizes of integers,
and often a few others, are supported by some programming languages. For
example, Java includes four signed integer sizes: **byte, short, int**, and **long**.
Some languages, for example, C++ and C#, include unsigned integer types,
which are types for integer values without signs. Unsigned types are often used
for binary data.

A signed integer value is represented in a computer by a string of bits, with
one of the bits (typically the leftmost) representing the sign. Most integer types
are supported directly by the hardware. One example of an integer type that is
not supported directly by the hardware is the long integer type of Python (F#
also provides such integers). Values of this type can have unlimited length. Long
integer values can be specified as literals, as in the following example:

```
243725839182756281923L
```

Integer arithmetic operations in Python that produce values too large to be
represented with **int** type store them as long integer type values.
A negative integer could be stored in sign-magnitude notation, in which the sign bit is set to indicate negative and the remainder of the bit string represents the absolute value of the number. Sign-magnitude notation, however, does not lend itself to computer arithmetic. Most computers now use a notation called two's complement to store negative integers, which is convenient for addition and subtraction. In two's-complement notation, the representation of a negative integer is formed by taking the logical complement of the positive version of the number and adding one. Ones-complement notation is still used by some computers. In ones-complement notation, the negative of an integer is stored as the logical complement of its absolute value. Ones-complement notation has the disadvantage that it has two representations of zero. See any book on assembly language programming for details of integer representations.

### 6.2.1.2 Floating-Point

**Floating-point** data types model real numbers, but the representations are only approximations for many real values. For example, neither of the fundamental numbers $\pi$ or $e$ (the base for the natural logarithms) can be correctly represented in floating-point notation. Of course, neither of these numbers can be precisely represented in any finite amount of computer memory. On most computers, floating-point numbers are stored in binary, which exacerbates the problem. For example, even the value 0.1 in decimal cannot be represented by a finite number of binary digits.\(^1\) Another problem with floating-point types is the loss of accuracy through arithmetic operations. For more information on the problems of floating-point notation, see any book on numerical analysis.

Floating-point values are represented as fractions and exponents, a form that is borrowed from scientific notation. Older computers used a variety of different representations for floating-point values. However, most newer machines use the IEEE Floating-Point Standard 754 format. Language implementors use whatever representation is supported by the hardware. Most languages include two floating-point types, often called *float* and *double*. The float type is the standard size, usually stored in four bytes of memory. The double type is provided for situations where larger fractional parts and/or a larger range of exponents is needed. Double-precision variables usually occupy twice as much storage as float variables and provide at least twice the number of bits of fraction.

The collection of values that can be represented by a floating-point type is defined in terms of precision and range. **Precision** is the accuracy of the fractional part of a value, measured as the number of bits. **Range** is a combination of the range of fractions and, more important, the range of exponents.

Figure 6.1 shows the IEEE Floating-Point Standard 754 format for single- and double-precision representation (IEEE, 1985). Details of the IEEE formats can be found in Tanenbaum (2005).

---

1. 0.1 in decimal is 0.0001100110011... in binary.
6.2.1.3 Complex

Some programming languages support a complex data type—for example, Fortran and Python. Complex values are represented as ordered pairs of floating-point values. In Python, the imaginary part of a complex literal is specified by following it with a $j$ or $J$—for example,

$$(7 + 3j)$$

Languages that support a complex type include operations for arithmetic on complex values.

6.2.1.4 Decimal

Most larger computers that are designed to support business systems applications have hardware support for decimal data types. Decimal data types store a fixed number of decimal digits, with the implied decimal point at a fixed position in the value. These are the primary data types for business data processing and are therefore essential to COBOL. C# and F# also have decimal data types.

Decimal types have the advantage of being able to precisely store decimal values, at least those within a restricted range, which cannot be done with floating-point. For example, the number 0.1 (in decimal) can be exactly represented in a decimal type, but not in a floating-point type, as is noted in Section 6.2.1.2. The disadvantages of decimal types are that the range of values is restricted because no exponents are allowed, and their representation in memory is mildly wasteful, for reasons discussed in the following paragraph.

Decimal types are stored very much like character strings, using binary codes for the decimal digits. These representations are called binary coded decimal (BCD). In some cases, they are stored one digit per byte, but in others, they are packed two digits per byte. Either way, they take more storage than binary representations. It takes at least four bits to code a decimal digit. Therefore, to store a six-digit coded decimal number requires 24 bits of memory.
However, it takes only 20 bits to store the same number in binary.\footnote{Of course, unless a program needs to maintain a large number of large decimal values, the difference is insignificant.} The operations on decimal values are done in hardware on machines that have such capabilities; otherwise, they are simulated in software.

### 6.2.2 Boolean Types

**Boolean** types are perhaps the simplest of all types. Their range of values has only two elements: one for true and one for false. They were introduced in ALGOL 60 and have been included in most general-purpose languages designed since 1960. One popular exception is C89, in which numeric expressions are used as conditionals. In such expressions, all operands with nonzero values are considered true, and zero is considered false. Although C99 and C++ have a Boolean type, they also allow numeric expressions to be used as if they were Boolean. This is not the case in the subsequent languages, Java and C#.

Boolean types are often used to represent switches or flags in programs. Although other types, such as integers, can be used for these purposes, the use of Boolean types is more readable.

A Boolean value could be represented by a single bit, but because a single bit of memory cannot be accessed efficiently on many machines, they are often stored in the smallest efficiently addressable cell of memory, typically a byte.

### 6.2.3 Character Types

Character data are stored in computers as numeric codings. Traditionally, the most commonly used coding was the 8-bit code ASCII (American Standard Code for Information Interchange), which uses the values 0 to 127 to code 128 different characters. ISO 8859-1 is another 8-bit character code, but it allows 256 different characters.

Because of the globalization of business and the need for computers to communicate with other computers around the world, the ASCII character set became inadequate. In response, in 1991, the Unicode Consortium published the UCS-2 standard, a 16-bit character set. This character code is often called Unicode. Unicode includes the characters from most of the world’s natural languages. For example, Unicode includes the Cyrillic alphabet, as used in Serbia, and the Thai digits. The first 128 characters of Unicode are identical to those of ASCII. Java was the first widely used language to use the Unicode character set. Since then, it has found its way into JavaScript, Python, Perl, C#, and F#.

After 1991, the Unicode Consortium, in cooperation with the International Standards Organization (ISO), developed a 4-byte character code named UCS-4, or UTF-32, which is described in the ISO/IEC 10646 Standard, published in 2000.
To provide the means of processing codings of single characters, most programming languages include a primitive type for them. However, Python supports single characters only as character strings of length 1.

6.3 Character String Types

A character string type is one in which the values consist of sequences of characters. Character string constants are used to label output, and the input and output of all kinds of data are often done in terms of strings. Of course, character strings also are an essential type for all programs that do character manipulation.

6.3.1 Design Issues

The two most important design issues that are specific to character string types are the following:

- Should strings be a special kind of character array or a primitive type?
- Should strings have static or dynamic length?

6.3.2 Strings and Their Operations

The most common string operations are assignment, catenation, substring reference, comparison, and pattern matching.

A substring reference is a reference to a substring of a given string. Substring references are discussed in the more general context of arrays, where the substring references are called slices.

In general, both assignment and comparison operations on character strings are complicated by the possibility of string operands of different lengths. For example, what happens when a longer string is assigned to a shorter string, or vice versa? Usually, simple and sensible choices are made for these situations, although programmers often have trouble remembering them.

In some languages, pattern matching is supported directly in the language. In others, it is provided by a function or class library.

If strings are not defined as a primitive type, string data is usually stored in arrays of single characters and referenced as such in the language. This is the approach taken by C and C++, which use char arrays to store character strings. These languages provide a collection of string operations through standard libraries. Many users of strings and many of the library functions use the convention that character strings are terminated with a special character, null, which is represented with zero. This is an alternative to maintaining the length of string variables. The library operations simply carry out their operations until the null character appears in the string being operated on. Library functions that produce strings often supply the null character. The character string
Character String Types

literals that are built by the compiler also have the null character. For example, consider the following declaration:

```c
char str[] = "apples";
```

In this example, `str` is an array of `char` elements, specifically `apples0`, where `0` is the null character.

Some of the most commonly used library functions for character strings in C and C++ are `strcpy`, which moves strings; `strcat`, which concatenates one given string onto another; `strcmp`, which lexicographically compares (by the order of their character codes) two given strings; and `strlen`, which returns the number of characters, not counting the null, in the given string. The parameters and return values for most of the string manipulation functions are `char` pointers that point to arrays of `char`. Parameters can also be string literals.

The string manipulation functions of the C standard library, which are also available in C++, are inherently unsafe and have led to numerous programming errors. The problem is that the functions in this library that move string data do not guard against overflowing the destination. For example, consider the following call to `strcpy`:

```c
strcpy(dest, src);
```

If the length of `dest` is 20 and the length of `src` is 50, `strcpy` will write over the 30 bytes that follow `dest`. The point is that `strcpy` does not know the length of `dest`, so it cannot ensure that the memory following it will not be overwritten. The same problem can occur with several of the other functions in the C string library. In addition to C-style strings, C++ also supports strings through its standard class library, which is also similar to that of Java. Because of the insecurities of the C string library, C++ programmers should use the `string` class from the standard library, rather than `char` arrays and the C string library.

In Java, strings are supported by the `String` class, whose values are constant strings, and the `StringBuffer` class, whose values are changeable and are more like arrays of single characters. These values are specified with methods of the `StringBuffer` class. C# and Ruby include string classes that are similar to those of Java.

Python includes strings as a primitive type and has operations for substring reference, catenation, indexing to access individual characters, as well as methods for searching and replacement. There is also an operation for character membership in a string. So, even though Python’s strings are primitive types, for character and substring references, they act very much like arrays of characters. However, Python strings are immutable, similar to the `String` class objects of Java.

In F#, strings are a class. Individual characters, which are represented in Unicode UTF-16, can be accessed, but not changed. Strings can be catenated with the `+` operator. In ML, string is a

**history note**

SNOBOL 4 was the first widely known language to support pattern matching.
primitive immutable type. It uses `^` for its catenation operator and includes functions for substring referencing and getting the size of a string.

Perl, JavaScript, Ruby, and PHP include built-in pattern-matching operations. In these languages, the pattern-matching expressions are somewhat loosely based on mathematical regular expressions. In fact, they are often called **regular expressions**. They evolved from the early UNIX line editor, `ed`, to become part of the UNIX shell languages. Eventually, they grew to their current complex form. There is at least one complete book on this kind of pattern-matching expressions (Friedl, 2006). In this section, we provide a brief look at the style of these expressions through two relatively simple examples.

Consider the following pattern expression:

```
/\[A-Za-z]\[A-Za-z\\d\]+/
```

This pattern matches (or describes) the typical name form in programming languages. The brackets enclose character classes. The first character class specifies all letters; the second specifies all letters and digits (a digit is specified with the abbreviation `\d`). If only the second character class were included, we could not prevent a name from beginning with a digit. The plus operator following the second category specifies that there must be one or more of what is in the category. So, the whole pattern matches strings that begin with a letter, followed by one or more letters or digits.

Next, consider the following pattern expression:

```
/\d+\.?\d*|\.\d+/
```

This pattern matches numeric literals. The `\.` specifies a literal decimal point. The question mark quantifies what it follows to have zero or one appearance. The vertical bar (`|`) separates two alternatives in the whole pattern. The first alternative matches strings of one or more digits, possibly followed by a decimal point, followed by zero or more digits; the second alternative matches strings that begin with a decimal point, followed by one or more digits.

Pattern-matching capabilities using regular expressions are included in the class libraries of C++, Java, Python, C#, and F#.

### 6.3.3 String Length Options

There are several design choices regarding the length of string values. First, the length can be static and set when the string is created. Such a string is called a **static length string**. This is the choice for the strings of Python, the immutable objects of Java’s `String` class, as well as similar classes in the C++ standard class library, Ruby’s built-in `String` class, and the .NET class library available to C# and F#.

---

3. The period must be “escaped” with the backslash because period has special meaning in a regular expression.
The second option is to allow strings to have varying length up to a declared and fixed maximum set by the variable’s definition, as exemplified by the strings in C and the C-style strings of C++. These are called **limited dynamic length strings**. Such string variables can store any number of characters between zero and the maximum. Recall that strings in C use a special character to indicate the end of the string’s characters, rather than maintaining the string length.

The third option is to allow strings to have varying length with no maximum, as in JavaScript, Perl, and the standard C++ library. These are called **dynamic length strings**. This option requires the overhead of dynamic storage allocation and deallocation but provides maximum flexibility.

### 6.3.4 Evaluation

String types are important to the writability of a language. Dealing with strings as arrays can be more cumbersome than dealing with a primitive string type. For example, consider a language that treats strings as arrays of characters and does not have a predefined function that does what `strcpy` in C does. Then, a simple assignment of one string to another would require a loop. The addition of strings as a primitive type to a language is not costly in terms of either language or compiler complexity. Therefore, it is difficult to justify the omission of primitive string types in some contemporary languages. Of course, providing strings through a standard library is nearly as convenient as having them as a primitive type.

String operations such as simple pattern matching and catenation are essential and should be included for string type values. Although dynamic length strings are obviously the most flexible, the overhead of their implementation must be weighed against that additional flexibility.

### 6.3.5 Implementation of Character String Types

Character string types could be supported directly in hardware; but in most cases, software is used to implement string storage, retrieval, and manipulation. When character string types are represented as character arrays, the language often supplies few operations.

A descriptor for a static character string type, which is required only during compilation, has three fields. The first field of every descriptor is the name of the type. In the case of static character strings, the second field is the type’s length (in characters). The third field is the address of the first character. This descriptor is shown in Figure 6.2. Limited dynamic strings require a run-time descriptor to store both the fixed maximum length and the current length, as shown in Figure 6.3. Dynamic length strings require a simpler run-time descriptor because only the current length needs to be stored. Although we depict descriptors as independent blocks of storage, in most cases, they are stored in the symbol table.
The limited dynamic strings of C and C++ do not require run-time descriptors, because the end of a string is marked with the null character. They do not need the maximum length, because index values in array references are not range checked in these languages.

Static length and limited dynamic length strings require no special dynamic storage allocation. In the case of limited dynamic length strings, sufficient storage for the maximum length is allocated when the string variable is bound to storage, so only a single allocation process is involved.

Dynamic length strings require more complex storage management. The length of a string, and therefore the storage to which it is bound, must grow and shrink dynamically.

There are three approaches to supporting the dynamic allocation and deallocation that is required for dynamic length strings. First, strings can be stored in a linked list, so that when a string grows, the newly required cells can come from anywhere in the heap. The drawbacks to this method are the extra storage occupied by the links in the list representation and the necessary complexity of string operations.

The second approach is to store strings as arrays of pointers to individual characters allocated in the heap. This method still uses extra memory, but string processing can be faster than with the linked-list approach.

The third alternative is to store complete strings in adjacent storage cells. The problem with this method arises when a string grows: How can storage that is adjacent to the existing cells continue to be allocated for the string variable? Frequently, such storage is not available. Instead, a new area of memory is found that can store the complete new string, and the old part is moved to this area. Then, the memory cells used for the old string are deallocated. This latter approach is the one typically used. The general problem of managing allocation and deallocation of variable-size segments is discussed in Section 6.11.7.3.

Although the linked-list method requires more storage, the associated allocation and deallocation processes are simple. However, some string operations
are slowed by the required pointer chasing. On the other hand, using adjacent memory for complete strings results in faster string operations and requires significantly less storage, but the allocation and deallocation processes are slower.

### 6.4 Enumeration Types

An **enumeration type** is one in which all of the possible values, which are named constants, are provided, or enumerated, in the definition. Enumeration types provide a way of defining and grouping collections of named constants, which are called **enumeration constants**. The definition of a typical enumeration type is shown in the following C# example:

```csharp
enum days {Mon, Tue, Wed, Thu, Fri, Sat, Sun};
```

The enumeration constants are typically implicitly assigned the integer values, 0, 1, . . . but can be explicitly assigned any integer literal in the type’s definition.

#### 6.4.1 Design Issues

The design issues for enumeration types are as follows:

- Is an enumeration constant allowed to appear in more than one type definition, and if so, how is the type of an occurrence of that constant in the program checked?
- Are enumeration values coerced to integer?
- Are any other types coerced to an enumeration type?

All of these design issues are related to type checking. If an enumeration variable is coerced to a numeric type, then there is little control over its range of legal operations or its range of values. If an `int` type value is coerced to an enumeration type, then an enumeration type variable could be assigned any integer value, whether it represented an enumeration constant or not.

#### 6.4.2 Designs

In languages that do not have enumeration types, programmers usually simulate them with integer values. For example, suppose we needed to represent colors in a C program and C did not have an enumeration type. We might use 0 to represent blue, 1 to represent red, and so forth. These values could be defined as follows:

```c
int red = 0, blue = 1;
```

Now, in the program, we could use `red` and `blue` as if they were of a color type. The problem with this approach is that because we have not defined a type for
our colors, there is no type checking when they are used. For example, it would be legal to add the two together, although that would rarely be an intended operation. They could also be combined with any other numeric type operand using any arithmetic operator, which would also rarely be useful. Furthermore, because they are just variables, they could be assigned any integer value, thereby destroying the relationship with the colors. This latter problem could be prevented by making them named constants.

C and Pascal were the first widely used languages to include an enumeration data type. C++ includes C’s enumeration types. In C++, we could have the following:

```cpp
enum colors {red, blue, green, yellow, black};
colors myColor = blue, yourColor = red;
```

The `colors` type uses the default internal values for the enumeration constants, 0, 1, ... , although the constants could have been assigned any integer literal (or any constant-valued expression). The enumeration values are coerced to `int` when they are put in integer context. This allows their use in any numeric expression. For example, if the current value of `myColor` is `blue`, then the expression

```cpp
myColor++
```

would assign `green` to `myColor`.

C++ also allows enumeration constants to be assigned to variables of any numeric type, though that would likely be an error. However, no other type value is coerced to an enumeration type in C++. For example,

```cpp
myColor = 4;
```

is illegal in C++. This assignment would be legal if the right side had been cast to `colors` type. This prevents some potential errors.

C++ enumeration constants can appear in only one enumeration type in the same referencing environment.

In 2004, an enumeration type was added to Java in Java 5.0. All enumeration types in Java are implicitly subclasses of the predefined class `Enum`. Because enumeration types are classes, they can have instance data fields, constructors, and methods. Syntactically, Java enumeration type definitions appear like those of C++, except that they can include fields, constructors, and methods. The possible values of an enumeration are the only possible instances of the class. All enumeration types inherit `toString`, as well as a few other methods. An array of the instances of an enumeration type can be fetched with the static method `values`. The internal numeric value of an enumeration variable can be fetched with the `ordinal` method. No expression of any other type can be assigned to an enumeration variable. Also, an enumeration variable is never coerced to any other type.
C# enumeration types are like those of C++, except that they are never coerced to integer. So, operations on enumeration types are restricted to those that make sense. Also, the range of values is restricted to that of the particular enumeration type.

In ML, enumeration types are defined as new types with `datatype` declarations. For example, we could have the following:

```ml
datatype weekdays = Monday | Tuesday | Wednesday | Thursday | Friday
```

The type of the elements of `weekdays` is integer.

F# has enumeration types that are similar to those of ML, except the reserved word `type` is used instead of `datatype` and the first value is preceded by an OR operator (`|`).

Interestingly, none of the relatively recent scripting languages include enumeration types. These include Perl, JavaScript, PHP, Python, Ruby, and Lua. Even Java was a decade old before enumeration types were added.

### 6.4.3 Evaluation

Enumeration types can provide advantages in both readability and reliability. Readability is enhanced very directly: Named values are easily recognized, whereas coded values are not.

In the area of reliability, the enumeration types of C#, F#, and Java 5.0 provide two advantages: (1) No arithmetic operations are legal on enumeration types; this prevents adding days of the week, for example, and (2) second, no enumeration variable can be assigned a value outside its defined range.4 If the `colors` enumeration type has 10 enumeration constants and uses `0..9` as its internal values, no number greater than 9 can be assigned to a `colors` type variable.

Because C treats enumeration variables like integer variables, it does not provide either of these two advantages.

C++ is a little better. Numeric values can be assigned to enumeration type variables only if they are cast to the type of the assigned variable. Numeric values assigned to enumeration type variables are checked to determine whether they are in the range of the internal values of the enumeration type. Unfortunately, if the user uses a wide range of explicitly assigned values, this checking is not effective. For example,

```c++
enum colors {red = 1, blue = 1000, green = 100000}
```

In this example, a value assigned to a variable of `colors` type will only be checked to determine whether it is in the range of `1..100000`.

---

4. In C# and F#, an integer value can be cast to an enumeration type and assigned to the name of an enumeration variable. Such values must be tested with `Enum.IsDefined` method before assigning them to the name of an enumeration variable.
6.5 Array Types

An array is a homogeneous aggregate of data elements in which an individual element is identified by its position in the aggregate, relative to the first element. The individual data elements of an array are of the same type. References to individual array elements are specified using subscript expressions. If any of the subscript expressions in a reference include variables, then the reference will require an additional run-time calculation to determine the address of the memory location being referenced.

In many languages, such as C, C++, Java, and C#, all of the elements of an array are required to be of the same type. In these languages, pointers and references are restricted to point to or reference a single type. So the objects or data values being pointed to or referenced are also of a single type. In some other languages, such as JavaScript, Python, and Ruby, variables are typeless references to objects or data values. In these cases, arrays still consist of elements of a single type, but the elements can reference objects or data values of different types. Such arrays are still homogeneous, because the array elements are of the same type.

C# and Java 5.0 provide generic arrays, that is, arrays whose elements are references to objects, through their class libraries. These are discussed in Section 6.5.3.

6.5.1 Design Issues

The primary design issues specific to arrays are the following:

- What types are legal for subscripts?
- Are subscripting expressions in element references range checked?
- When are subscript ranges bound?
- When does array allocation take place?
- Are ragged or rectangular multidimensioned arrays allowed, or both?
- Can arrays be initialized when they have their storage allocated?
- What kinds of slices are allowed, if any?

In the following sections, examples of the design choices made for the arrays of the most common programming languages are discussed.

6.5.2 Arrays and Indices

Specific elements of an array are referenced by means of a two-level syntactic mechanism, where the first part is the aggregate name, and the second part is a possibly dynamic selector consisting of one or more items known as subscripts or indices. If all of the subscripts in a reference are constants, the selector is static; otherwise, it is dynamic. The selection operation can be thought of as a mapping from the array name and the set of subscript values to an element in
the aggregate. Indeed, arrays are sometimes called finite mappings. Symbolically, this mapping can be shown as

\[
\text{array}_\text{name}(\text{subscript}_\text{value}_\text{list}) \rightarrow \text{element}
\]

The syntax of array references is fairly universal: The array name is followed by the list of subscripts, which is surrounded by either parentheses or brackets. In some languages that provide multidimensioned arrays as arrays of arrays, each subscript appears in its own brackets. A problem with using parentheses to enclose subscript expressions is that they often are also used to enclose the parameters in subprogram calls; this use makes references to arrays appear exactly like those calls. For example, consider the following Ada assignment statement:

\[
\text{Sum} := \text{Sum} + \text{B}(\text{I});
\]

Because parentheses are used for both subprogram parameters and array subscripts in Ada, both program readers and compilers are forced to use other information to determine whether \( \text{B}(\text{I}) \) in this assignment is a function call or a reference to an array element. This results in reduced readability.

The designers of Ada specifically chose parentheses to enclose subscripts so there would be uniformity between array references and function calls in expressions, in spite of potential readability problems. They made this choice in part because both array element references and function calls are mappings. Array element references map the subscripts to a particular element of the array. Function calls map the actual parameters to the function definition and, eventually, a functional value.

Most languages other than Fortran and Ada use brackets to delimit their array indices.

Two distinct types are involved in an array type: the element type and the type of the subscripts. The type of the subscripts is often integer.

Early programming languages did not specify that subscript ranges must be implicitly checked. Range errors in subscripts are common in programs, so requiring range checking is an important factor in the reliability of languages. Many contemporary languages also do not specify range checking of subscripts, but Java, ML, and C# do.

Subscripting in Perl is a bit unusual in that although the names of all arrays begin with at signs (@), because array elements are always scalars and the names of scalars always begin with dollar signs ($), references to array elements use dollar signs rather than at signs in their names. For example, for the array @list, the second element is referenced with @list[1].

One can reference an array element in Perl with a negative subscript, in which case the subscript value is an offset from the end of the array. For example, if the array @list has five elements with the subscripts 0..4, $list[-2]
references the element with the subscript 3. A reference to a nonexistent element in Perl yields `undef`, but no error is reported.

### 6.5.3 Subscript Bindings and Array Categories

The binding of the subscript type to an array variable is usually static, but the subscript value ranges are sometimes dynamically bound.

In some languages, the lower bound of the subscript range is implicit. For example, in the C-based languages, the lower bound of all subscript ranges is fixed at 0. In some other languages, the lower bounds of the subscript ranges must be specified by the programmer.

There are four categories of arrays, based on the binding to subscript ranges, the binding to storage, and from where the storage is allocated. The category names indicate the design choices of these three. In the first three of these categories, once the subscript ranges are bound and the storage is allocated, they remain fixed for the lifetime of the variable. Of course, when the subscript ranges are fixed, the array cannot change size.

A **static array** is one in which the subscript ranges are statically bound and storage allocation is static (done before run time). The advantage of static arrays is efficiency: No dynamic allocation or deallocation is required. The disadvantage is that the storage for the array is fixed for the entire execution time of the program.

A **fixed stack-dynamic array** is one in which the subscript ranges are statically bound, but the allocation is done at declaration elaboration time during execution. The advantage of fixed stack-dynamic arrays over static arrays is space efficiency. A large array in one subprogram can use the same space as a large array in a different subprogram, as long as both subprograms are not active at the same time. The same is true if the two arrays are in different blocks that are not active at the same time. The disadvantage is the required allocation and deallocation time.

A **fixed heap-dynamic array** is similar to a fixed stack-dynamic array, in that the subscript ranges and the storage binding are both fixed after storage is allocated. The differences are that both the subscript ranges and storage bindings are done when the user program requests them during execution, and the storage is allocated from the heap, rather than the stack. The advantage of fixed heap-dynamic arrays is flexibility—the array’s size always fits the problem. The disadvantage is allocation time from the heap, which is longer than allocation time from the stack.

A **heap-dynamic array** is one in which the binding of subscript ranges and storage allocation is dynamic and can change any number of times during the array’s lifetime. The advantage of heap-dynamic arrays over the others is flexibility: Arrays can grow and shrink during program execution as the need for space changes. The disadvantage is that allocation and deallocation take longer and may happen many times during execution of the program. Examples of the four categories are given in the following paragraphs.

Arrays declared in C and C++ functions that include the `static` modifier are static.
Arrays that are declared in C and C++ functions without the `static` specifier are examples of fixed stack-dynamic arrays.

C and C++ also provide fixed heap-dynamic arrays. The standard C library functions `malloc` and `free`, which are general heap allocation and deallocation operations, respectively, can be used for C arrays. C++ uses the operators `new` and `delete` to manage heap storage. An array is treated as a pointer to a collection of storage cells, where the pointer can be indexed, as discussed in Section 6.11.5.

In Java, all non-generic arrays are fixed heap-dynamic. Once created, these arrays keep the same subscript ranges and storage. C# also provides fixed heap-dynamic arrays.

Objects of the C# `List` class are generic heap-dynamic arrays. These array objects are created without any elements, as in

```java
List<String> stringList = new List<String>();
```

Elements are added to this object with the `Add` method, as in

```java
stringList.Add("Michael");
```

Access to elements of these arrays is through subscripting.

Java includes a generic class similar to C#’s `List`, named `ArrayList`. It is different from C#’s `List` in that subscripting is not supported—`get` and `set` methods must be used to access the elements.

A Perl array can be made to grow by using the `push` (puts one or more new elements on the end of the array) and `unshift` (puts one or more new elements on the beginning of the array), or by assigning a value to the array specifying a subscript beyond the highest current subscript of the array. An array can be made to shrink to no elements by assigning it the empty list, `()`.

The length of an array is defined to be the largest subscript plus one.

Like Perl, JavaScript allows arrays to grow with the `push` and `unshift` methods and shrink by setting them to the empty list. However, negative subscripts are not supported.

JavaScript arrays can be sparse, meaning the subscript values need not be contiguous. For example, suppose we have an array named `list` that has 10 elements with the subscripts `0..9`. Consider the following assignment statement:

```javascript
list[50] = 42;
```

Now, `list` has 11 elements and length 51. The elements with subscripts `11..49` are not defined and therefore do not require storage. A reference to a nonexistent element in a JavaScript array yields `undefined`.

Arrays in Python, Ruby, and Lua can be made to grow only through methods to add elements or catenate other arrays. Ruby and Lua support negative

---

5. The subscript range could just as easily have been `1000 .. 1009`. 
subscripts, but Python does not. In Python, Ruby, and Lua an element or slice of an array can be deleted. A reference to a nonexistent element in Python results in a run-time error, whereas a similar reference in Ruby and Lua yields \texttt{nil} and no error is reported.

Although the ML definition does not include arrays, its widely used implementation, SML/NJ, does.

The only predefined collection type that is part of F# is the array (other collection types are provided through the .NET Framework Library). These arrays are like those of C#. A \texttt{foreach} statement is included in the language for array processing.

### 6.5.4 Array Initialization

Some languages provide the means to initialize arrays at the time their storage is allocated. C, C++, Java, and C# allow initialization of their arrays. Consider the following C declaration:

```c
int list [] = {4, 5, 7, 83};
```

The array \texttt{list} is created and initialized with the values 4, 5, 7, and 83. The compiler also sets the length of the array. This is meant to be a convenience but is not without cost. It effectively removes the possibility that the system could detect some kinds of programmer errors, such as mistakenly leaving a value out of the list.

As discussed in Section 6.3.2, character strings in C and C++ are implemented as arrays of \texttt{char}. These arrays can be initialized to string constants, as in

```c
char name [] = "freddie";
```

The array \texttt{name} will have eight elements, because all strings are terminated with a null character (zero), which is implicitly supplied by the system for string constants.

Arrays of strings in C and C++ can also be initialized with string literals. For example,

```c
char *names [] = {"Bob", "Jake", "Darcie"};
```

This example illustrates the nature of character literals in C and C++. In the previous example of a string literal being used to initialize the \texttt{char} array \texttt{name}, the literal is taken to be a \texttt{char} array. But in the latter example (\texttt{names}), the literals are taken to be pointers to characters, so the array is an array of pointers to characters. For example, \texttt{names[0]} is a pointer to the letter 'B' in the literal character array that contains the characters 'B', 'o', 'b', and the null character.

In Java, similar syntax is used to define and initialize an array of references to \texttt{String} objects. For example,

```java
String[] names = ["Bob", "Jake", "Darcie"];
```
6.5.5 Array Operations

An array operation is one that operates on an array as a unit. The most common array operations are assignment, catenation, comparison for equality and inequality, and slices, which are discussed separately in Section 6.5.5.

The C-based languages do not provide any array operations, except through the methods of Java, C++, and C#. Perl supports array assignments but does not support comparisons.

Python’s arrays are called lists, although they have all the characteristics of dynamic arrays. Because the objects can be of any types, these arrays are heterogeneous. Python provides array assignment, although it is only a reference change. Python also has operations for array catenation (+) and element membership (in). It includes two different comparison operators: one that determines whether the two variables reference the same object (is) and one that compares all corresponding objects in the referenced objects, regardless of how deeply they are nested, for equality (==).

Like Python, the elements of Ruby’s arrays are references to objects. And like Python, when a == operator is used between two arrays, the result is true only if the two arrays have the same length and the corresponding elements are equal. Ruby’s arrays can be catenated with an Array method.

F# includes many array operators in its Array module. Among these are Array.append, Array.copy, and Array.length.

Arrays and their operations are the heart of APL; it is the most powerful array-processing language ever devised. Because of its relative obscurity and its lack of effect on subsequent languages, however, we present here only a glimpse into its array operations.

In APL, the four basic arithmetic operations are defined for vectors (single-dimensioned arrays) and matrices, as well as scalar operands. For example,

\[ A + B \]

is a valid expression, whether \( A \) and \( B \) are scalar variables, vectors, or matrices.

APL includes a collection of unary operators for vectors and matrices, some of which are as follows (where \( V \) is a vector and \( M \) is a matrix):

\[
\begin{align*}
\phi V & \text{ reverses the elements of } V \\
\phi M & \text{ reverses the columns of } M \\
\theta M & \text{ reverses the rows of } M \\
\otimes M & \text{ transposes } M \text{ (its rows become its columns and vice versa)} \\
\div M & \text{ inverts } M 
\end{align*}
\]

APL also includes several special operators that take other operators as operands. One of these is the inner product operator, which is specified with a period (\(.\)). It takes two operands, which are binary operators. For example,

\[ +.\times \]
is a new operator that takes two arguments, either vectors or matrices. It first multiplies the corresponding elements of two arguments, and then it sums the results. For example, if \( \mathbf{A} \) and \( \mathbf{B} \) are vectors,

\[ \mathbf{A} \times \mathbf{B} \]

is the mathematical inner product of \( \mathbf{A} \) and \( \mathbf{B} \) (a vector of the products of the corresponding elements of \( \mathbf{A} \) and \( \mathbf{B} \)). The statement

\[ \mathbf{A} + \times \mathbf{B} \]

is the sum of the inner product of \( \mathbf{A} \) and \( \mathbf{B} \). If \( \mathbf{A} \) and \( \mathbf{B} \) are matrices, this expression specifies the matrix multiplication of \( \mathbf{A} \) and \( \mathbf{B} \).

The special operators of APL are actually functional forms, which are described in Chapter 15.

### 6.5.6 Rectangular and Jagged Arrays

A **rectangular array** is a multidimensioned array in which all of the rows have the same number of elements and all of the columns have the same number of elements. Rectangular arrays model rectangular tables exactly.

A **jagged array** is one in which the lengths of the rows need not be the same. For example, a jagged matrix may consist of three rows, one with 5 elements, one with 7 elements, and one with 12 elements. This also applies to the columns and higher dimensions. So, if there is a third dimension (layers), each layer can have a different number of elements. Jagged arrays are made possible when multidimensioned arrays are actually arrays of arrays. For example, a matrix would appear as an array of single-dimensioned arrays.

C, C++, and Java support jagged arrays but not rectangular arrays. In those languages, a reference to an element of a multidimensioned array uses a separate pair of brackets for each dimension. For example,

\[ \text{myArray}[3][7] \]

C# and F# support both rectangular and jagged arrays. For rectangular arrays, all subscript expressions in references to elements are placed in a single pair of brackets. For example,

\[ \text{myArray}[3, 7] \]

### 6.5.7 Slices

A **slice** of an array is some substructure of that array. For example, if \( \mathbf{A} \) is a matrix, then the first row of \( \mathbf{A} \) is one possible slice, as are the last row and the first column. It is important to realize that a slice is not a new data type. Rather, it is a mechanism for referencing part of an array as a unit. If arrays cannot be manipulated as units in a language, that language has no use for slices.
Consider the following Python declarations:

```python
vector = [2, 4, 6, 8, 10, 12, 14, 16]
mat = [[1, 2, 3], [4, 5, 6], [7, 8, 9]]
```

Recall that the default lower bound for Python arrays is 0. The syntax of a Python slice reference is a pair of numeric expressions separated by a colon. The first is the first subscript of the slice; the second is the first subscript after the last subscript in the slice. Therefore, `vector[3:6]` is a three-element array with the fourth through sixth elements of `vector` (those elements with the subscripts 3, 4, and 5). A row of a matrix is specified by giving just one subscript. For example, `mat[1]` refers to the second row of `mat`; a part of a row can be specified with the same syntax as a part of a single-dimensioned array. For example, `mat[0][0:2]` refers to the first and second element of the first row of `mat`, which is `[1, 2]`.

Python also supports more complex slices of arrays. For example, `vector[0:7:2]` references every other element of `vector`, up to but not including the element with the subscript 7, starting with the subscript 0, which is `[2, 6, 10, 14]`.

Perl supports slices of two forms, a list of specific subscripts or a range of subscripts. For example,

```perl
@list[1..5] = @list2[3, 5, 7, 9, 13];
```

Notice that slice references use array names, not scalar names, because slices are arrays (not scalars).

Ruby supports slices with the `slice` method of its `Array` object, which can take three forms of parameters. A single integer expression parameter is interpreted as a subscript, in which case `slice` returns the element with the given subscript. If `slice` is given two integer expression parameters, the first is interpreted as a beginning subscript and the second is interpreted as the number of elements in the slice. For example, suppose `list` is defined as follows:

```ruby
list = [2, 4, 6, 8, 10]
```

`list.slice(2, 2)` returns `[6, 8]`. The third parameter form for `slice` is a range, which has the form of an integer expression, two periods, and a second integer expression. With a range parameter, `slice` returns an array of the element with the given range of subscripts. For example, `list.slice(1..3)` returns `[4, 6, 8]`.

### 6.5.8 Evaluation

Arrays have been included in virtually all programming languages. The primary advances since their introduction in Fortran I have been slices and dynamic arrays. As discussed in Section 6.6, the latest advances in arrays have been in associative arrays.
### 6.5.9 Implementation of Array Types

Implementing arrays requires considerably more compile-time effort than does implementing primitive types. The code to allow accessing of array elements must be generated at compile time. At run time, this code must be executed to produce element addresses. There is no way to precompute the address to be accessed by a reference such as

\[ \text{list}[k] \]

A single-dimensioned array is implemented as a list of adjacent memory cells. Suppose the array list is defined to have a subscript range lower bound of 0. The access function for list is often of the form

\[ \text{address(list}[k]] = \text{address(list}[0]] + k \times \text{element_size} \]

where the first operand of the addition is the constant part of the access function, and the second is the variable part.

If the element type is statically bound and the array is statically bound to storage, then the value of the constant part can be computed before run time. However, the addition and multiplication operations must be done at run time.

The generalization of this access function for an arbitrary lower bound is

\[ \text{address(list}[k]] = \text{address(list}[\text{lower_bound}]] + ((k - \text{lower_bound}) \times \text{element_size}) \]

The compile-time descriptor for single-dimensioned arrays can have the form shown in Figure 6.4. The descriptor includes information required to construct the access function. If run-time checking of index ranges is not done and the attributes are all static, then only the access function is required during execution; no descriptor is needed. If run-time checking of index ranges is done, then those index ranges may need to be stored in a run-time descriptor. If the subscript ranges of a particular array type are static, then the ranges may be incorporated into the code that does the checking, thus eliminating the need for the run-time descriptor. If any of the descriptor entries are dynamically bound, then those parts of the descriptor must be maintained at run time.

**Figure 6.4**

<table>
<thead>
<tr>
<th>Array</th>
<th>Element type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Index type</td>
<td>Index lower bound</td>
</tr>
<tr>
<td>Index upper bound</td>
<td>Address</td>
</tr>
</tbody>
</table>
True multidimensional arrays, that is, those that are not arrays of arrays, are more complex to implement than single-dimensioned arrays, although the extension to more dimensions is straightforward. Hardware memory is linear—just a simple sequence of bytes. So values of data types that have two or more dimensions must be mapped onto the single-dimensioned memory. There are two ways in which multidimensional arrays can be mapped to one dimension: row major order and column major order (not used in any widely used language). In **row major order**, the elements of the array that have as their first subscript the lower bound value of that subscript are stored first, followed by the elements of the second value of the first subscript, and so forth. If the array is a matrix, it is stored by rows. For example, if the matrix had the values

\[
\begin{array}{ccc}
3 & 4 & 7 \\
6 & 2 & 5 \\
1 & 3 & 8 \\
\end{array}
\]

it would be stored in row major order as

\[
3, 4, 7, 6, 2, 5, 1, 3, 8
\]

The access function for a multidimensional array is the mapping of its base address and a set of index values to the address in memory of the element specified by the index values. The access function for two-dimensional arrays stored in row major order can be developed as follows. In general, the address of an element is the base address of the structure plus the element size times the number of elements that precede it in the structure. For a matrix in row major order, the number of elements that precede an element is the number of rows above the element times the size of a row, plus the number of elements to the left of the element in its row. This is illustrated in Figure 6.5, in which we assume that subscript lower bounds are all zero.

To get an actual address value, the number of elements that precede the desired element must be multiplied by the element size. Now, the access function can be written as

![Figure 6.5](image-url)
location(a[i, j]) = address of a[0, 0] 
+ (((number of rows above the \textit{i}th row) * (size of a row)) 
+ (number of elements left of the \textit{j}th column)) * 
 element size)

Because the number of rows above the \textit{i}th row is \textit{i} and the number of elements 
to the left of the \textit{j}th column is \textit{j}, we have

location(a[i, j]) = address of a[0, 0] + ((\textit{i} * n) + \textit{j}) * 
 element size)

where \textit{n} is the number of elements per row. The first term is the constant part 
and the last is the variable part.

The generalization to arbitrary lower bounds results in the following access 
function:

location(a[i, j]) = address of a[\text{row}_{lb}, \text{col}_{lb}] 
= (((\textit{i} - \text{row}_{lb}) * \textit{n}) + (\textit{j} - \text{col}_{lb})) * \text{element size}

where \text{row}_{lb} is the lower bound of the rows and \text{col}_{lb} is the lower bound of 
the columns. This can be rearranged to the form

location(a[i, j]) = address of a[\text{row}_{lb}, \text{col}_{lb}] 
- (((\text{row}_{lb} * n) + \text{col}_{lb}) * \text{element size}) 
+ (((\textit{i} * n) + \textit{j}) * \text{element size})

where the first two terms are the constant part and the last is the variable part. 
This can be generalized rather easily to an arbitrary number of dimensions.

For each dimension of an array, one add and one multiply instruction are 
required for the access function. Therefore, accesses to elements of arrays with 
several subscripts are costly. The compile-time descriptor for a multidimensional 
array is shown in Figure 6.6.

**Figure 6.6**
A compile-time 
descriptor for a 
multidimensional array
6.6 Associative Arrays

An **associative array** is an unordered collection of data elements that are indexed by an equal number of values called **keys**. In the case of non-associative arrays, the indices never need to be stored (because of their regularity). In an associative array, however, the user-defined keys must be stored in the structure. So each element of an associative array is in fact a pair of entities, a key and a value. We use Perl’s design of associative arrays to illustrate this data structure. Associative arrays are also supported directly by Python, Ruby, and Lua and by the standard class libraries of Java, C++, C#, and F#.

The only design issue that is specific for associative arrays is the form of references to their elements.

6.6.1 Structure and Operations

In Perl, associative arrays are called **hashes**, because in the implementation their elements are stored and retrieved with hash functions. The namespace for Perl hashes is distinct: Every hash variable name must begin with a percent sign (%). Each hash element consists of two parts: a key, which is a string, and a value, which is a scalar (number, string, or reference). Hashes can be set to literal values with the assignment statement, as in

```perl
%salaries = ("Gary" => 75000, "Perry" => 57000, "Mary" => 55750, "Cedric" => 47850);
```

Individual element values are referenced using notation that is similar to that used for Perl arrays. The key value is placed in braces and the hash name is replaced by a scalar variable name that is the same except for the first character. Although hashes are not scalars, the value parts of hash elements are scalars, so references to hash element values use scalar names. Recall that scalar variable names begin with dollar signs ($). So, an assignment of 58850 to the element of `%salaries` with the key "Perry" would appear as follows:

```perl
$salaries{"Perry"} = 58850;
```

A new element is added using the same assignment statement form. An element can be removed from the hash with the `delete` operator, as in the following:

```perl
delete $salaries{"Gary"};
```

The entire hash can be emptied by assigning the empty literal to it, as in the following:

```perl
@salaries = ();
```

The size of a Perl hash is dynamic: It grows when an element is added and shrinks when an element is deleted, and also when it is emptied by assignment.
How and where did you first become involved with computing? Before I entered college in 1978, I had no idea about computing. I remember that I tried to read a book on programming in Fortran but did not pass the initial chapter on definitions for variables and constants.

In my first year in college I took a Programming 101 course in Fortran. At that time we ran our programming assignments in an IBM 370 mainframe. We had to punch cards with our code, surround the deck with some fixed JCL cards and give it to an operator. Some time later (often a few hours) we got a listing with the results, which frequently were only compiler errors.

Soon after that a friend of mine brought from abroad a microcomputer, a Z80 CPU with 4K bytes of memory. We started to do all kinds of programs for this machine, all in assembly—or, more exactly, in machine code, as it did not have an assembler. We wrote our programs in assembly, then translated them by hand to hexadecimal to enter them into memory to run.

Since then I was hooked.

There have been few successful programming languages designed in academic environments in the last 25 years. Although you are an academic, Lua was designed for very practical applications. Do you consider Lua an academic or an industrial language? Lua is certainly an industrial language, but with an academic “accent.” Lua was created for two industrial applications, and it has been used in industrial applications all its life. We tried to be very pragmatic on its design. However, except for its first version, we were never under the typical pressure from an industrial environment. We always had the luxury of choosing when to release a new version or of choosing whether to accept user demands. That gave us some latitude that other languages have not enjoyed.

More recently, we have done some academic research with Lua. But it is a long process to merge these academic results into the official distribution; more often than not these results have little direct impact on Lua. Nevertheless, there have been some nice exceptions, such as the register-based virtual machine and “ephemerons” (to appear in Lua 5.2).

You have said Lua was raised, rather than designed. Can you comment on what you meant and what you think are the benefits of this approach? We meant that most important pieces of Lua were not present in its first version. The language started as a very small and simple language and got several of its relevant features as it evolved.

Before talking about the benefits (and the drawbacks) of this approach, let me make it clear that we did not choose that approach. We never thought, “let us grow a new language.” It just happened.

I guess that a most difficult part when designing a language is to foresee how different mechanisms will interact in daily use. By raising a language—that is, creating it piece by piece—you may avoid most of those interaction problems, as you can think about each new feature only after the rest of the language is in place and has been tested by real users in real applications.

Of course, this approach has a major drawback, too: You may arrive at a point where a most-needed new feature is incompatible with what you already have in place.

Lua has changed in a variety of ways since it was first released in 1994. You have said that there have been times when you regretted not including a Boolean type in Lua. Why didn’t you simply add one? This may sound funny, but what we really missed was the value “false”; we had no use for a “true” value.
Like the original Lisp, Lua treated \texttt{nil} as false and everything else as true. The problem is that \texttt{nil} also represents an uninitialized variable. There was no way to distinguish between an uninitialized variable from a false variable. So, we needed a false value, to make that distinction possible. But the true value was useless; a 1 or any other constant was good enough.

I guess this is a typical example where our “industrial” mind conflicted with our “academic” mind. A really pragmatic mind would add the Boolean type without thinking twice. But our academic mind was upset by this inelegance. In the end the pragmatic side won, but it took some time.

**What were the most important Lua features, other than the preprocessor, that later became recognized as misfeatures and were removed from the language?** I do not remember other big misfeatures. We did remove several features from Lua, but mostly because they were superseded by a new, usually “better” in some sense, feature. This happened with tag methods (superseded by metamethods), weak references in the C API (superseded by weak tables), and upvalues (superseded by proper lexical scoping).

**When a new feature for Lua that would break backward compatibility is considered, how is that decision made?** These are always hard decisions. First, we try to find some other format that could avoid or at least reduce the incompatibility. If that is not possible, we try to provide easy ways around the incompatibility. (For instance, if we remove a function from the core library we may provide a separated implementation that the programmer may incorporate into her code.) Also, we try to measure how difficult it will be to detect and correct the incompatibility. If the new feature creates syntax errors (e.g., a new reserved word), that is not that bad; we may even provide an automatic tool to fix old code. However, if the new feature may produce subtle bugs (e.g., a preexisting function returning a different result), we consider it unacceptable.

**Were iterator methods, like those of Ruby, considered for Lua, rather than the \texttt{for} statement that was added? What considerations led to the choice?** They were not only considered, they were actually implemented! Since version 3.1 (from 1998), Lua has had a function “\texttt{foreach}”, that applies a given function to all pairs in a table. Similarly, with “\texttt{gsub}” it is easy to apply a given function to each character in a string.

Instead of a special “block” mechanism for the iterator body, Lua has used first-class functions for the task. See the next example:

```lua
--'t' is a table from names to values
--the next "loop" prints all keys with values greater than 10
foreach(t, function(key, value)
    if value > 10 then print(key) end
end)
```

However, when we first implemented iterators, functions in Lua did not have full lexical scoping. Moreover, the syntax is a little heavy (macros would help). Also, exit statements (break and return) are always confusing when used inside iteration bodies. So, in the end we decided for the \texttt{for} statement.

But “true iterators” are still a useful design in Lua, even more now that functions have proper lexical scoping. In my Lua book, I end the chapter about the \texttt{for} statement with a discussion of true iterators.

**Can you briefly describe what you mean when you describe Lua as an extensible extension language?** It is an “extensible language” because it is easy to register new functions and types defined in other languages. So it is easy to extend the language. From a more concrete point of view, it is easy to call C from Lua.

It is an “extension language” because it is easy to use Lua to extend an application, to morph Lua into a macro language for the application. (This is “scripting” in its purer meaning.) From a more concrete point of view, it is easy to call Lua from C.

**Data structures have evolved from arrays, records, and hashes to combinations of these. Can you estimate the significance of Lua’s tables in the evolution of data structures in programming languages?** I do not think the Lua table has had any significance in the evolution of other languages. Maybe that will change in the future, but I am not sure about it. In my view, the main benefit offered by Lua tables is its simplicity, an “all-in-one” solution. But this simplicity has its costs: For instance, static analysis of Lua programs is very hard, partially because of tables being so generic and ubiquitous. Each language has its own priorities.
of the empty literal. The `exists` operator returns true or false, depending on whether its operand key is an element in the hash. For example,

```
if (exists $salaries{"Shelly"}) . . .
```

The `keys` operator, when applied to a hash, returns an array of the keys of the hash. The `values` operator does the same for the values of the hash. The `each` operator iterates over the element pairs of a hash.

Python’s associative arrays, which are called `dictionaries`, are similar to those of Perl, except the values are all references to objects. The associative arrays supported by Ruby are similar to those of Python, except that the keys can be any object,6 rather than just strings. There is a progression from Perl’s hashes, in which the keys must be strings, to PHP’s arrays, in which the keys can be integers or strings, to Ruby’s hashes, in which any type object can be a key.

PHP’s arrays are both normal arrays and associative arrays. They can be treated as either. The language provides functions that allow both indexed and hashed access to elements. An array can have elements that are created with simple numeric indices and elements that are created with string hash keys.

In Lua, the table type is the only data structure. A Lua table is an associative array in which both the keys and the values can be any type. A table can be used as a traditional array, an associative array, or a record (struct). When used as a traditional array or an associative array, brackets are used around the keys. When used as a record, the keys are the field names and references to fields can use dot notation (record_name.field_name).

The use of Lua’s associative arrays as records is discussed in Section 6.7.

C# and F# support associative arrays through a .NET class.

An associative array is much better than an array if searches of the elements are required, because the implicit hashing operation used to access elements is very efficient. Furthermore, associative arrays are ideal when the data to be stored is paired, as with employee names and their salaries. On the other hand, if every element of a list must be processed, it is more efficient to use an array.

### 6.6.2 Implementing Associative Arrays

The implementation of Perl’s associative arrays is optimized for fast lookups, but it also provides relatively fast reorganization when array growth requires it. A 32-bit hash value is computed for each entry and is stored with the entry, although an associative array initially uses only a small part of the hash value. When an associative array must be expanded beyond its initial size, the hash function need not be changed; rather, more bits of the hash value are used. Only

---

6. Objects that change do not make good keys, because the changes could change the hash function value. Therefore, arrays and hashes are never used as keys.
half of the entries must be moved when this happens. So, although expansion of an associative array is not free, it is not as costly as might be expected.

The elements in PHP’s arrays are placed in memory through a hash function. However, all elements are linked together in the order in which they were created. The links are used to support iterative access to elements through the current and next functions.

6.7 Record Types

A record is an aggregate of data elements in which the individual elements are identified by names and accessed through offsets from the beginning of the structure.

There is frequently a need in programs to model a collection of data in which the individual elements are not of the same type or size. For example, information about a college student might include name, student number, grade point average, and so forth. A data type for such a collection might use a character string for the name, an integer for the student number, a floating-point for the grade point average, and so forth. Records are designed for this kind of need.

It may appear that records and heterogeneous arrays are the same, but that is not the case. The elements of a heterogeneous array are all references to data objects that reside in scattered locations, often on the heap. The elements of a record are of potentially different sizes and reside in adjacent memory locations.

Records have been part of all of the most popular programming languages, except pre-90 versions of Fortran, since the early 1960s, when they were introduced by COBOL. In some languages that support object-oriented programming, data classes serve as records.

In C, C++, and C#, records are supported with the struct data type. In C++, structures are a minor variation on classes. In C#, structs also are related to classes, but are quite different from them. C# structs are stack-allocated value types, as opposed to class objects, which are heap-allocated reference types. Structs in C++ and C# are normally used as encapsulation structures, rather than data structures. They are further discussed in this capacity in Chapter 11. Structs are also included in ML and F#.

In Python and Ruby, records can be implemented as hashes, which themselves can be elements of arrays.

The following sections describe how records are declared or defined, how references to fields within records are made, and the common record operations.

The following design issues are specific to records:

- What is the syntactic form of references to fields?
- Are elliptical references allowed?
6.7.1 Definitions of Records

The fundamental difference between a record and an array is that record elements, or fields, are not referenced by indices. Instead, the fields are named with identifiers, and references to the fields are made using these identifiers. Another difference between arrays and records is that records in some languages are allowed to include unions, which are discussed in Section 6.10.

The COBOL form of a record declaration, which is part of the data division of a COBOL program, is illustrated in the following example:

```
01 EMPLOYEE-RECORD.
  02 EMPLOYEE-NAME.
    05 FIRST PICTURE IS X(20).
    05 Middle PICTURE IS X(10).
    05 LAST  PICTURE IS X(20).
  02 HOURLY-RATE  PICTURE IS 99V99.
```

The EMPLOYEE-RECORD record consists of the EMPLOYEE-NAME record and the HOURLY-RATE field. The numerals 01, 02, and 05 that begin the lines of the record declaration are level numbers, which indicate by their relative values the hierarchical structure of the record. Any line that is followed by a line with a higher-level number is itself a record. The PICTURE clauses show the formats of the field storage locations, with X(20) specifying 20 alphanumeric characters and 99V99 specifying four decimal digits with the decimal point in the middle.

In Java and C#, records can be defined as data classes, with nested records defined as nested classes. Data members of such classes serve as the record fields.

As stated previously, Lua’s associative arrays can be conveniently used as records. For example, consider the following declaration:

```
employee.name = "Freddie"
employee.hourlyRate = 13.20
```

These assignment statements create a table (record) named employee with two elements (fields) named name and hourlyRate, both initialized.

6.7.2 References to Record Fields

References to the individual fields of records are syntactically specified by several different methods, two of which name the desired field and its enclosing records. COBOL field references have the form

```
field_name OF record_name_1 OF . . . OF record_name_n
```

where the first record named is the smallest or innermost record that contains the field. The next record name in the sequence is that of the record that
contains the previous record, and so forth. For example, the `Middle` field in the COBOL record example above can be referenced with

```
Middle OF EMPLOYEE-NAME OF EMPLOYEE-RECORD
```

Most of the other languages use **dot notation** for field references, where the components of the reference are connected with periods. Names in dot notation have the opposite order of COBOL references: They use the name of the largest enclosing record first and the field name last. For example, if `Middle` is a field in the `Employee_Name` record which is embedded in the `Employee_Record` record, it would be referenced with the following:

```
Employee_Record.Employee_Name.Middle
```

References to elements in a Lua table can appear in the syntax of record field references, as seen in the assignment statements in Section 6.7.1. Such references could also have the form of normal table elements—for example, `employee["name"]`.

A **fully qualified reference** to a record field is one in which all intermediate record names, from the largest enclosing record to the specific field, are named in the reference. In the COBOL example above the field reference is fully qualified. As an alternative to fully qualified references, COBOL allows **elliptical references** to record fields. In an elliptical reference, the field is named, but any or all of the enclosing record names can be omitted, as long as the resulting reference is unambiguous in the referencing environment. For example, `FIRST`, `FIRST OF Employee_Name`, and `FIRST OF Employee_Record` are elliptical references to the employee's first name in the COBOL record declared above. Although elliptical references are a programmer convenience, they require a compiler to have elaborate data structures and procedures in order to correctly identify the referenced field. They are also somewhat detrimental to readability.

### 6.7.3 Evaluation

Records are frequently valuable data types in programming languages. The design of record types is straightforward, and their use is safe.

Records and arrays are closely related structural forms, and therefore it is interesting to compare them. Arrays are used when all the data values have the same type and/or are processed in the same way. This processing is easily done when there is a systematic way of sequencing through the structure. Such processing is well supported by using dynamic subscripting as the addressing method.

Records are used when the collection of data values is heterogeneous and the different fields are not processed in the same way. Also, the fields of a record often need not be processed in a particular order. Field names are like literal, or
constant, subscripts. Because they are static, they provide very efficient access to the fields. Dynamic subscripts could be used to access record fields, but it would disallow type checking and would also be slower.

Records and arrays represent thoughtful and efficient methods of fulfilling two separate but related applications of data structures.

### 6.7.4 Implementation of Record Types

The fields of records are stored in adjacent memory locations. But because the sizes of the fields are not necessarily the same, the access method used for arrays is not used for records. Instead, the offset address, relative to the beginning of the record, is associated with each field. Field accesses are all handled using these offsets. The compile-time descriptor for a record has the general form shown in Figure 6.7. Run-time descriptors for records are unnecessary.

### 6.8 Tuple Types

A tuple is a data type that is similar to a record, except that the elements are not named.

Python includes an immutable tuple type. If a tuple needs to be changed, it can be converted to an array with the `list` function. After the change, it can be converted back to a tuple with the `tuple` function. One use of tuples is when an array must be write protected, such as when it is sent as a parameter to an external function and the user does not want the function to be able to modify the parameter.
Python's tuples are closely related to its lists, except that tuples are immutable. A tuple is created by assigning a tuple literal, as in the following example:

```python
myTuple = (3, 5.8, 'apple')
```

Notice that the elements of a tuple need not be of the same type.

The elements of a tuple can be referenced with indexing in brackets, as in the following:

```python
myTuple[1]
```

This references the first element of the tuple, because tuple indexing begins at 1.

Tuples can be catenated with the plus (+) operator. They can be deleted with the `del` statement. There are also other operators and functions that operate on tuples.

ML includes a tuple data type. An ML tuple must have at least two elements, whereas Python's tuples can be empty or contain one element. As in Python, an ML tuple can include elements of mixed types. The following statement creates a tuple:

```ml
val myTuple = (3, 5.8, 'apple');
```

The syntax of a tuple element access is as follows:

```ml
#1(myTuple);
```

This references the first element of the tuple.

A new tuple type can be defined in ML with a type declaration, such as the following:

```ml
type intReal = int * real;
```

Values of this type consist of an integer and a real.

F# also has tuples. A tuple is created by assigning a tuple value, which is a list of expressions separated by commas and delimited by parentheses, to a name in a `let` statement. If a tuple has two elements, they can be referenced with the functions `fst` and `snd`, respectively. The elements of a tuple with more than two elements are often referenced with a tuple pattern on the left side of a `let` statement. A tuple pattern is simply a sequence of names, one for each element of the tuple, with or without the delimiting parentheses. When a tuple pattern is the left side of a `let` construct, it is a multiple assignment. For example, consider the following `let` constructs:

```fsharp
let tup = (3, 5, 7);
let a, b, c = tup;
```

This assigns 3 to `a`, 5 to `b`, and 7 to `c`.

Tuples are used in Python, ML, and F# to allow functions to return multiple values.
6.9 List Types

Lists were first supported in the first functional programming language, Lisp. They have always been part of the functional languages, but in recent years they have found their way into some imperative languages.

Lists in Scheme and Common Lisp are delimited by parentheses and the elements are not separated by any punctuation. For example,

\[(A \ B \ C \ D)\]

Nested lists have the same form, so we could have

\[(A \ (B \ C) \ D)\]

In this list, \((B \ C)\) is a list nested inside the outer list.

Data and code have the same syntactic form in Lisp and its descendants. If the list \((A \ B \ C)\) is interpreted as code, it is a call to the function \(A\) with parameters \(B\) and \(C\).

The fundamental list operations in Scheme are two functions that take lists apart and two that build lists. The \(\text{CAR}\) function returns the first element of its list parameter. For example, consider the following example:

\[(\text{CAR} \ ' (A \ B \ C))\]

The quote before the parameter list is to prevent the interpreter from considering the list a call to the \(A\) function with the parameters \(B\) and \(C\), in which case it would interpret it. This call to \(\text{CAR}\) returns \(A\).

The \(\text{CDR}\) function returns its parameter list minus its first element. For example, consider the following example:

\[(\text{CDR} \ ' (A \ B \ C))\]

This function call returns the list \((B \ C)\).

Common Lisp also has the functions \(\text{FIRST}\) (same as \(\text{CAR}\)), \(\text{SECOND}\), \ldots, \(\text{TENTH}\), which return the element of their list parameters that is specified by their names.

In Scheme and Common Lisp, new lists are constructed with the \(\text{CONS}\) and \(\text{LIST}\) functions. The function \(\text{CONS}\) takes two parameters and returns a new list with its first parameter as the first element and its second parameter as the remainder of that list. For example, consider the following:

\[(\text{CONS} \ 'A \ ' (B \ C))\]

This call returns the new list \((A \ B \ C)\).

The \(\text{LIST}\) function takes any number of parameters and returns a new list with the parameters as its elements. For example, consider the following call to \(\text{LIST}\):

\[(\text{LIST} \ 'A \ 'B \ ' (C \ D))\]
This call returns the new list \((A \ B \ (C \ D))\).

ML has lists and list operations, although their appearance is not like those of Scheme. Lists are specified in square brackets, with the elements separated by commas, as in the following list of integers:

\[ [5, 7, 9] \]

[] is the empty list, which could also be specified with nil.

The Scheme \texttt{CONS} function is implemented as a binary infix operator in ML, represented as ::. For example,

\[ 3 :: [5, 7, 9] \]

returns the following new list: \([3, 5, 7, 9]\).

The elements of a list must be of the same type, so the following list would be illegal:

\[ [5, 7.3, 9] \]

ML has functions that correspond to Scheme’s \texttt{CAR} and \texttt{CDR}, named \texttt{hd} (head) and \texttt{tl} (tail). For example,

\[ \text{hd} [5, 7, 9] \text{ is } 5 \]

\[ \text{tl} [5, 7, 9] \text{ is } [7, 9] \]

Lists and list operations in Scheme and ML are more fully discussed in Chapter 15.

Lists in F# are related to those of ML with a few notable differences. Elements of a list in F# are separated by semicolons, rather than the commas of ML. The operations \texttt{hd} and \texttt{tl} are the same, but they are called as methods of the \texttt{List} class, as in \texttt{List.hd [1; 3; 5; 7]}, which returns 1. The \texttt{CONS} operation of F# is specified as two colons, as in ML.

Python includes a list data type, which also serves as Python’s arrays. Unlike the lists of Scheme, Common Lisp, ML, and F#, the lists of Python are mutable. They can contain any data value or object. A Python list is created with an assignment of a list value to a name. A list value is a sequence of expressions that are separated by commas and delimited with brackets. For example, consider the following statement:

\[ \text{myList} = [3, 5.8, "grape"] \]

The elements of a list are referenced with subscripts in brackets, as in the following example:

\[ x = \text{myList}[1] \]

This statement assigns 5.8 to \(x\). The elements of a list are indexed starting at zero. List elements also can be updated by assignment. A list element can be deleted with \texttt{del}, as in the following statement:

\[ \text{del myList[1]} \]
This statement removes the second element of myList.

Python includes a powerful mechanism for creating arrays called list comprehensions. A list comprehension is an idea derived from set notation. It first appeared in the functional programming language Haskell (see Chapter 15). The mechanics of a list comprehension is that a function is applied to each of the elements of a given array and a new array is constructed from the results. The syntax of a Python list comprehension is as follows:

\[
\text{[expression for iterate_var in array if condition]}
\]

Consider the following example:

\[
[x * x for x in range(12) if x % 3 == 0]
\]

The range function creates the array \([0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12]\). The conditional filters out all numbers in the array that are not evenly divisible by 3. Then, the expression squares the remaining numbers. The results of the squaring are collected in an array, which is returned. This list comprehension returns the following array:

\([0, 9, 36, 81]\)

Slices of lists are also supported in Python.

Haskell’s list comprehensions have the following form:

\([\text{body | qualifiers}]\)

For example, consider the following definition of a list:

\([n * n | n <- [1..10]]\)

This defines a list of the squares of the numbers from 1 to 10.

F# includes list comprehensions, which in that language can also be used to create arrays. For example, consider the following statement:

\(\text{let myArray = [|for i in 1 .. 5 -> (i * i) |];;}\)

This statement creates the array \([1; 4; 9; 16; 25]\) and names it myArray.

Recall from Section 6.5 that C# and Java support generic heap-dynamic collection classes, List and ArrayList, respectively. These structures are actually lists.

### 6.10 Union Types

A union is a type whose variables may store different type values at different times during program execution. As an example of the need for a union type, consider a table of constants for a compiler, which is used to store the constants
found in a program being compiled. One field of each table entry is for the value of the constant. Suppose that for a particular language being compiled, the types of constants were integer, floating-point, and Boolean. In terms of table management, it would be convenient if the same location, a table field, could store a value of any of these three types. Then all constant values could be addressed in the same way. The type of such a location is, in a sense, the union of the three value types it can store.

### 6.10.1 Design Issues

The problem of type checking union types, which is discussed in Section 6.12, is their major design issue.

### 6.10.2 Discriminated Versus Free Unions

C and C++ provide union constructs in which there is no language support for type checking. In C and C++, the `union` construct is used to specify union structures. The unions in these languages are called free unions, because programmers are allowed complete freedom from type checking in their use. For example, consider the following C union:

```c
union flexType {
    int intEl;
    float floatEl;
};
union flexType el1;
float x;
...
el1.intEl = 27;
x = el1.floatEl;
```

This last assignment is not type checked, because the system cannot determine the current type of the current value of `el1`, so it assigns the bit string representation of 27 to the `float` variable `x`, which of course is nonsense.

Type checking of unions requires that each union construct include a type indicator. Such an indicator is called a tag, or discriminant, and a union with a discriminant is called a discriminated union. The first language to provide discriminated unions was ALGOL 68. They are now supported by ML, Haskell, and F#.

### 6.10.3 Unions in F#

A union is declared in F# with a type statement using OR operators (|) to define the components. For example, we could have the following:

```fsharp
type intReal =
    | IntValue of int
    | RealValue of float;;
```
In this example, `intReal` is the union type. `IntValue` and `RealValue` are constructors. Values of type `intReal` can be created using the constructors as if they were a function, as in the following examples:

```fsharp
let ir1 = IntValue 17;;
let ir2 = RealValue 3.4;;
```

Accessing the value of a union is done with a pattern-matching structure. Pattern matching in F# is specified with the `match` reserved word. The general form of the construct is as follows:

```
match pattern with
| expression_list1 -> expression1
| . . .
| expression_listn -> expressionn
```

The pattern can be any data type. The expression list can include wild card characters (`_`) or be solely a wild card character. For example, consider the following match construct:

```fsharp
let a = 7;;
let b = "grape";;
let x = match (a, b) with
| 4, "apple" -> apple
| _, "grape" -> grape
| _ -> fruit;;
```

To display the type of the `intReal` union, the following function could be used:

```fsharp
let printType value =
    match value with
    | IntValue value -> printfn "It is an integer"
    | RealValue value -> printfn "It is a float";;
```

The following lines show calls to this function and the output:

```
printType ir1;;
It is an integer
printType ir2;;
It is a float
```

---

7. The `let` statement is used to assign values to names and to create a static scope; the double semicolons are used to terminate statements when the F# interactive interpreter is being used.
6.10.4 Evaluation

Unions are potentially unsafe constructs in some languages. They are one of the reasons why C and C++ are not strongly typed: These languages do not allow type checking of references to their unions. On the other hand, unions can be safely used, as in their design in ML, Haskell, and F#.

Neither Java nor C# includes unions, which may be reflective of the growing concern for safety in some programming languages.

6.10.5 Implementation of Union Types

Unions are implemented by simply using the same address for every possible variant. Sufficient storage for the largest variant is allocated.

6.11 Pointer and Reference Types

A pointer type is one in which the variables have a range of values that consists of memory addresses and a special value, nil. The value nil is not a valid address and is used to indicate that a pointer cannot currently be used to reference a memory cell.

Pointers are designed for two distinct kinds of uses. First, pointers provide some of the power of indirect addressing, which is frequently used in assembly language programming. Second, pointers provide a way to manage dynamic storage. A pointer can be used to access a location in an area where storage is dynamically allocated called a heap.

Variables that are dynamically allocated from the heap are called heap-dynamic variables. They often do not have identifiers associated with them and thus can be referenced only by pointer or reference type variables. Variables without names are called anonymous variables. It is in this latter application area of pointers that the most important design issues arise.

Pointers, unlike arrays and records, are not structured types, although they are defined using a type operator (* in C and C++). Furthermore, they are also different from scalar variables because they are used to reference some other variable, rather than being used to store data. These two categories of variables are called reference types and value types, respectively.

Both kinds of uses of pointers add writability to a language. For example, suppose it is necessary to implement a dynamic structure like a binary tree in a language that does not have pointers or dynamic storage. This would require the programmer to provide and maintain a pool of available tree nodes, which would probably be implemented in parallel arrays. Also, it would be necessary for the programmer to guess the maximum number of required nodes. This is clearly an awkward and error-prone way to deal with binary trees.

Reference variables, which are discussed in Section 6.11.6, are closely related to pointers.
6.11.1 Design Issues

The primary design issues particular to pointers are the following:

- What are the scope and lifetime of a pointer variable?
- What is the lifetime of a heap-dynamic variable (the value a pointer references)?
- Are pointers restricted as to the type of value to which they can point?
- Are pointers used for dynamic storage management, indirect addressing, or both?
- Should the language support pointer types, reference types, or both?

6.11.2 Pointer Operations

Languages that provide a pointer type usually include two fundamental pointer operations: assignment and dereferencing. The first operation sets a pointer variable's value to some useful address. If pointer variables are used only to manage dynamic storage, then the allocation mechanism, whether by operator or built-in subprogram, serves to initialize the pointer variable. If pointers are used for indirect addressing to variables that are not heap dynamic, then there must be an explicit operator or built-in subprogram for fetching the address of a variable, which can then be assigned to the pointer variable.

An occurrence of a pointer variable in an expression can be interpreted in two distinct ways. First, it could be interpreted as a reference to the contents of the memory cell to which it is bound, which in the case of a pointer is an address. This is exactly how a nonpointer variable in an expression would be interpreted, although in that case its value likely would not be an address.

However, the pointer also could be interpreted as a reference to the value in the memory cell pointed to by the memory cell to which the pointer variable is bound. In this case, the pointer is interpreted as an indirect reference. The former case is a normal pointer reference; the latter is the result of dereferencing the pointer. Dereferencing, which takes a reference through one level of indirection, is the second fundamental pointer operation.

Dereferencing of pointers can be either explicit or implicit. In many contemporary languages, it occurs only when explicitly specified. In C++, it is explicitly specified with the asterisk (*) as a prefix unary operator. Consider the following example of dereferencing: If \( \text{ptr} \) is a pointer variable with the value 7080 and the cell whose address is 7080 has the value 206, then the assignment \( j = *\text{ptr} \) sets \( j \) to 206. This process is shown in Figure 6.8.

When pointers point to records, the syntax of the references to the fields of these records varies among languages. In C and C++, there are two ways a pointer to a record can be used to reference a field in that record. If a pointer variable \( p \) points to a record with a field named \( \text{age} \), \( (*p) . \text{age} \) can be used to
refer to that field. The operator \(-\rightarrow\), when used between a pointer to a struct and a field of that struct, combines dereferencing and field reference. For example, the expression \(p \rightarrow age\) is equivalent to \((*p).age\).

Languages that provide pointers for the management of a heap must include an explicit allocation operation. Allocation is sometimes specified with a subprogram, such as `malloc` in C. In languages that support object-oriented programming, allocation of heap objects is often specified with the `new` operator. C++, which does not provide implicit deallocation, uses `delete` as its deallocation operator.

### 6.11.3 Pointer Problems

The first high-level programming language to include pointer variables was PL/I, in which pointers could be used to refer to both heap-dynamic variables and other program variables. The pointers of PL/I were highly flexible, but their use could lead to several kinds of programming errors. Some of the problems of PL/I pointers are also present in the pointers of subsequent languages. Some recent languages, such as Java, have replaced pointers completely with reference types, which, along with implicit deallocation, minimize the primary problems with pointers. A reference type is really only a pointer with restricted operations.

#### 6.11.3.1 Dangling Pointers

A **dangling pointer**, or **dangling reference**, is a pointer that contains the address of a heap-dynamic variable that has been deallocated. Dangling pointers are dangerous for several reasons. First, the location being pointed to may have been reallocated to some new heap-dynamic variable. If the new variable is not the same type as the old one, type checks of uses of the dangling pointer are invalid. Even if the new dynamic variable is the same type, its new value will have no relationship to the old pointer’s dereferenced value. Furthermore, if the dangling pointer is used to change the heap-dynamic variable, the value of the new heap-dynamic variable will be destroyed. Finally, it is possible that
the location now is being temporarily used by the storage management system, possibly as a pointer in a chain of available blocks of storage, thereby allowing a change to the location to cause the storage manager to fail.

The following sequence of operations creates a dangling pointer in many languages:

1. A new heap-dynamic variable is created and pointer \( p_1 \) is set to point at it.
2. Pointer \( p_2 \) is assigned \( p_1 \)'s value.
3. The heap-dynamic variable pointed to by \( p_1 \) is explicitly deallocated (possibly setting \( p_1 \) to \( \text{nil} \)), but \( p_2 \) is not changed by the operation. \( p_2 \) is now a dangling pointer. If the deallocation operation did not change \( p_1 \), both \( p_1 \) and \( p_2 \) would be dangling. (Of course, this is a problem of aliasing—\( p_1 \) and \( p_2 \) are aliases.)

For example, in C++ we could have the following:

```cpp
int * arrayPtr1;
int * arrayPtr2 = new int[100];
arrayPtr1 = arrayPtr2;
delete [] arrayPtr2;
// Now, arrayPtr1 is dangling, because the heap storage // to which it was pointing has been deallocated.
```

In C++, both \( \text{arrayPtr1} \) and \( \text{arrayPtr2} \) are now dangling pointers, because the C++ \text{delete} \ operator has no effect on the value of its operand pointer. In C++, it is common (and safe) to follow a \text{delete} \ operator with an assignment of zero, which represents null, to the pointer whose pointed-to value has been deallocated.

Notice that the explicit deallocation of dynamic variables is the cause of dangling pointers.

### 6.11.3.2 Lost Heap-Dynamic Variables

A lost heap-dynamic variable is an allocated heap-dynamic variable that is no longer accessible to the user program. Such variables are often called \text{garbage}, because they are not useful for their original purpose, and they also cannot be reallocated for some new use in the program. Lost heap-dynamic variables are most often created by the following sequence of operations:

1. Pointer \( p_1 \) is set to point to a newly created heap-dynamic variable.
2. \( p_1 \) is later set to point to another newly created heap-dynamic variable.

The first heap-dynamic variable is now inaccessible, or lost. This is sometimes called \text{memory leakage}. Memory leakage is
a problem, regardless of whether the language uses implicit or explicit deallocation. In the following sections, we investigate how language designers have dealt with the problems of dangling pointers and lost heap-dynamic variables.

6.11.4 Pointers in C and C++

In C and C++, pointers can be used in the same ways as addresses are used in assembly languages. This means they are extremely flexible but must be used with great care. This design offers no solutions to the dangling pointer or lost heap-dynamic variable problems. However, the fact that pointer arithmetic is possible in C and C++ makes their pointers more interesting than those of the other programming languages.

C and C++ pointers can point at any variable, regardless of where it is allocated. In fact, they can point anywhere in memory, whether there is a variable there or not, which is one of the dangers of such pointers.

In C and C++, the asterisk (*) denotes the dereferencing operation and the ampersand (&) denotes the operator for producing the address of a variable. For example, consider the following code:

```c
int *ptr;
int count, init;
...
ptr = &init;
count = *ptr;
```

The assignment to the variable `ptr` sets it to the address of `init`. The assignment to `count` dereferences `ptr` to produce the value at `init`, which is then assigned to `count`. So, the effect of the two assignment statements is to assign the value of `init` to `count`. Notice that the declaration of a pointer specifies its domain type.

Pointers can be assigned the address value of any variable of the correct domain type, or they can be assigned the constant zero, which is used for `nil`.

Pointer arithmetic is also possible in some restricted forms. For example, if `ptr` is a pointer variable that is declared to point at some variable of some data type, then

```c
ptr + index
```

is a legal expression. The semantics of such an expression is as follows. Instead of simply adding the value of `index` to `ptr`, the value of `index` is first scaled by the size of the memory cell (in memory units) to which `ptr` is pointing (its base type). For example, if `ptr` points to a memory cell for a type that is four memory units in size, then `index` is multiplied by 4, and the result is added to `ptr`. The primary purpose of this sort of address arithmetic is array manipulation. The following discussion is related to single-dimensional arrays only.
In C and C++, all arrays use zero as the lower bound of their subscript ranges, and array names without subscripts always refer to the address of the first element. Consider the following declarations:

```c
int list [10];
int *ptr;
```

Now consider the assignment

```c
ptr = list;
```

This assigns the address of `list[0]` to `ptr`. Given this assignment, the following are true:

- *(ptr + 1)* is equivalent to `list[1].`
- *(ptr + index)* is equivalent to `list[index].`
- `ptr[index]` is equivalent to `list[index].`

It is clear from these statements that the pointer operations include the same scaling that is used in indexing operations. Furthermore, pointers to arrays can be indexed as if they were array names.

Pointers in C and C++ can point to functions. This feature is used to pass functions as parameters to other functions. Pointers are also used for parameter passing, as discussed in Chapter 9.

C and C++ include pointers of type `void *`, which can point at values of any type. In effect they are generic pointers. However, type checking is not a problem with `void *` pointers, because these languages disallow dereferencing them. One common use of `void *` pointers is as the types of parameters of functions that operate on memory. For example, suppose we wanted a function to move a sequence of bytes of data from one place in memory to another. It would be most general if it could be passed two pointers of any type. This would be legal if the corresponding formal parameters in the function were `void *` type. The function could then convert them to `char *` type and do the operation, regardless of what type pointers were sent as actual parameters.

### 6.11.5 Reference Types

A reference type variable is similar to a pointer, with one important and fundamental difference: A pointer refers to an address in memory, while a reference refers to an object or a value in memory. As a result, although it is natural to perform arithmetic on addresses, it is not sensible to do arithmetic on references.

C++ includes a special kind of reference type that is used primarily for the formal parameters in function definitions. A C++ reference type variable is a constant pointer that is always implicitly dereferenced. Because a C++ reference type variable is a constant, it must be initialized with the address of some
variable in its definition, and after initialization a reference type variable can never be set to reference any other variable. The implicit dereference prevents assignment to the address value of a reference variable.

Reference type variables are specified in definitions by preceding their names with ampersands (\&). For example,

```cpp
int result = 0;
int &ref_result = result;
... 
ref_result = 100;
```

In this code segment, `result` and `ref_result` are aliases.

When used as formal parameters in function definitions, C++ reference types provide for two-way communication between the caller function and the called function. This is not possible with nonpointer primitive parameter types, because C++ parameters are passed by value. Passing a pointer as a parameter accomplishes the same two-way communication, but pointer formal parameters require explicit dereferencing, making the code less readable and less safe. Reference parameters are referenced in the called function exactly as are other parameters. The calling function need not specify that a parameter whose corresponding formal parameter is a reference type is anything unusual. The compiler passes addresses, rather than values, to reference parameters.

In their quest for increased safety over C++, the designers of Java removed C++-style pointers altogether. Unlike C++ reference variables, Java reference variables can be assigned to refer to different class instances; they are not constants. All Java class instances are referenced by reference variables. That is, in fact, the only use of reference variables in Java. These issues are further discussed in Chapter 12.

In the following, `String` is a standard Java class:

```java
String str1;
... 
str1 = "This is a Java literal string";
```

In this code, `str1` is defined to be a reference to a `String` class instance or object. It is initially set to null. The subsequent assignment sets `str1` to reference the `String object, "This is a Java literal string".

Because Java class instances are implicitly deallocated (there is no explicit deallocation operator), there cannot be dangling references in Java.

C# includes both the references of Java and the pointers of C++. However, the use of pointers is strongly discouraged. In fact, any subprogram that uses pointers must include the `unsafe` modifier. Note that although objects pointed to by references are implicitly deallocated, that is not true for objects pointed to by pointers. Pointers were included in C# primarily to allow C# programs to interoperate with C and C++ code.

All variables in the object-oriented languages Smalltalk, Python, Ruby, and Lua are references. They are always implicitly dereferenced. Furthermore, the direct values of these variables cannot be accessed.
6.11.6 Evaluation

The problems of dangling pointers and garbage have already been discussed at length. The problems of heap management are discussed in Section 6.11.7.3.

Pointers have been compared with the goto. The goto statement widens the range of statements that can be executed next. Pointer variables widen the range of memory cells that can be referenced by a variable. Perhaps the most damning statement about pointers was made by Hoare (1973): “Their introduction into high-level languages has been a step backward from which we may never recover.”

On the other hand, pointers are essential in some kinds of programming applications. For example, pointers are necessary to write device drivers, in which specific absolute addresses must be accessed.

The references of Java and C# provide some of the flexibility and the capabilities of pointers, without the hazards. It remains to be seen whether programmers will be willing to trade the full power of C and C++ pointers for the greater safety of references. The extent to which C# programs use pointers will be one measure of this.

6.11.7 Implementation of Pointer and Reference Types

In most languages, pointers are used in heap management. The same is true for Java and C# references, as well as the variables in Smalltalk and Ruby, so we cannot treat pointers and references separately. First, we briefly describe how pointers and references are represented internally. We then discuss two possible solutions to the dangling pointer problem. Finally, we describe the major problems with heap-management techniques.

6.11.7.1 Representations of Pointers and References

In most larger computers, pointers and references are single values stored in memory cells. However, in early microcomputers based on Intel microprocessors, addresses have two parts: a segment and an offset. So, pointers and references are implemented in these systems as pairs of 16-bit cells, one for each of the two parts of an address.

6.11.7.2 Solutions to the Dangling-Pointer Problem

There have been several proposed solutions to the dangling-pointer problem. Among these are tombstones (Lomet, 1975), in which every heap-dynamic variable includes a special cell, called a tombstone, that is itself a pointer to the heap-dynamic variable. The actual pointer variable points only at tombstones and never to heap-dynamic variables. When a heap-dynamic variable is deallocated, the tombstone remains but is set to nil, indicating that the heap-dynamic variable no longer exists. This approach prevents a pointer from ever pointing to a deallocated variable. Any reference to any pointer that points to a nil tombstone can be detected as an error.
Tombstones are costly in both time and space. Because tombstones are never deallocated, their storage is never reclaimed. Every access to a heap-dynamic variable through a tombstone requires one more level of indirection, which requires an additional machine cycle on most computers. Apparently none of the designers of the more popular languages have found the additional safety to be worth this additional cost, because no widely used language uses tombstones.

An alternative to tombstones is the locks-and-keys approach used in the implementation of UW-Pascal (Fischer and LeBlanc, 1977, 1980). In this compiler, pointer values are represented as ordered pairs (key, address), where the key is an integer value. Heap-dynamic variables are represented as the storage for the variable plus a header cell that stores an integer lock value. When a heap-dynamic variable is allocated, a lock value is created and placed both in the lock cell of the heap-dynamic variable and in the key cell of the pointer that is specified in the call to \texttt{new}. Every access to the dereferenced pointer compares the key value of the pointer to the lock value in the heap-dynamic variable. If they match, the access is legal; otherwise the access is treated as a run-time error. Any copies of the pointer value to other pointers must copy the key value. Therefore, any number of pointers can reference a given heap-dynamic variable. When a heap-dynamic variable is deallocated with \texttt{dispose}, its lock value is cleared to an illegal lock value. Then, if a pointer other than the one specified in the \texttt{dispose} is dereferenced, its address value will still be intact, but its key value will no longer match the lock, so the access will not be allowed.

Of course, the best solution to the dangling-pointer problem is to take deallocation of heap-dynamic variables out of the hands of programmers. If programs cannot explicitly deallocate heap-dynamic variables, there will be no dangling pointers. To do this, the run-time system must implicitly deallocate heap-dynamic variables when they are no longer useful. Lisp systems have always done this. Both Java and C# also use this approach for their reference variables. Recall that C#s pointers do not include implicit deallocation.

\section*{6.11.7.3 Heap Management}

Heap management can be a very complex run-time process. We examine the process in two separate situations: one in which all heap storage is allocated and deallocated in units of a single size, and one in which variable-size segments are allocated and deallocated. Note that for deallocation, we discuss only implicit approaches. Our discussion will be brief and far from comprehensive, since a thorough analysis of these processes and their associated problems is not so much a language design issue as it is an implementation issue.

\textbf{Single-Size Cells} The simplest situation is when all allocation and deallocation is of single-size cells. It is further simplified when every cell already contains a pointer. This is the scenario of many implementations of Lisp, where the problems of dynamic storage allocation were first encountered on a large scale. All Lisp programs and most Lisp data consist of cells in linked lists.
In a single-size allocation heap, all available cells are linked together using the pointers in the cells, forming a list of available space. Allocation is a simple matter of taking the required number of cells from this list when they are needed. Deallocation is a much more complex process. A heap-dynamic variable can be pointed to by more than one pointer, making it difficult to determine when the variable is no longer useful to the program. Simply because one pointer is disconnected from a cell obviously does not make it garbage; there could be several other pointers still pointing to the cell.

In Lisp, several of the most frequent operations in programs create collections of cells that are no longer accessible to the program and therefore should be deallocated (put back on the list of available space). One of the fundamental design goals of Lisp was to ensure that reclamation of unused cells would not be the task of the programmer but rather that of the run-time system. This goal left Lisp implementors with the fundamental design question: When should deallocation be performed?

There are several different approaches to garbage collection. The two most common traditional techniques are in some ways opposite processes. These are named reference counters, in which reclamation is incremental and is done when inaccessible cells are created, and mark-sweep, in which reclamation occurs only when the list of available space becomes empty. These two methods are sometimes called the **eager approach** and the **lazy approach**, respectively. Many variations of these two approaches have been developed. In this section, however, we discuss only the basic processes.

The reference counter method of storage reclamation accomplishes its goal by maintaining in every cell a counter that stores the number of pointers that are currently pointing at the cell. Embedded in the decrement operation for the reference counters, which occurs when a pointer is disconnected from the cell, is a check for a zero value. If the reference counter reaches zero, it means that no program pointers are pointing at the cell, and it has thus become garbage and can be returned to the list of available space.

There are three distinct problems with the reference counter method. First, if storage cells are relatively small, the space required for the counters is significant. Second, some execution time is obviously required to maintain the counter values. Every time a pointer value is changed, the cell to which it was pointing must have its counter decremented, and the cell to which it is now pointing must have its counter incremented. In a language like Lisp, in which nearly every action involves changing pointers, that can be a significant portion of the total execution time of a program. Of course, if pointer changes are not too frequent, this is not a problem. Some of the inefficiency of reference counters can be eliminated by an approach named **deferred reference counting**, which avoids reference counters for some pointers. The third problem is that complications arise when a collection of cells is connected circularly. The problem here is that each cell in the circular list has a reference counter value of at least 1, which prevents it from being collected and placed back on the list of available space. A solution to this problem can be found in Friedman and Wise (1979).
The advantage of the reference counter approach is that it is intrinsically incremental. Its actions are interleaved with those of the application, so it never causes significant delays in the execution of the application.

The original mark-sweep process of garbage collection operates as follows: The run-time system allocates storage cells as requested and disconnects pointers from cells as necessary, without regard for storage reclamation (allowing garbage to accumulate), until it has allocated all available cells. At this point, a mark-sweep process is begun to gather all the garbage left floating around in the heap. To facilitate the process, every heap cell has an extra indicator bit or field that is used by the collection algorithm.

The mark-sweep process consists of three distinct phases. First, all cells in the heap have their indicators set to indicate they are garbage. This is, of course, a correct assumption for only some of the cells. The second part, called the marking phase, is the most difficult. Every pointer in the program is traced into the heap, and all reachable cells are marked as not being garbage. After this, the third phase, called the sweep phase, is executed: All cells in the heap that have not been specifically marked as still being used are returned to the list of available space.

To illustrate the flavor of algorithms used to mark the cells that are currently in use, we provide the following simple version of a marking algorithm. We assume that all heap-dynamic variables, or heap cells, consist of an information part; a part for the mark, named marker; and two pointers named llink and rlink. These cells are used to build directed graphs with at most two edges leading from any node. The marking algorithm traverses all spanning trees of the graphs, marking all cells that are found. Like other graph traversals, the marking algorithm uses recursion.

\[
\text{for every pointer } r \text{ do}
\]
\[
\text{mark}(r)
\]

\[
\text{void mark}(\text{void } * \text{ ptr}) \{
\text{if } (\text{ptr} \neq 0)
\text{if } (*\text{ptr.marker is not marked}) \{
\text{set } *\text{ptr.marker}
\text{mark}(\text{ptr.llink})
\text{mark}(\text{ptr.rlink})
\}
\}
\]

An example of the actions of this procedure on a given graph is shown in Figure 6.9. This simple marking algorithm requires a great deal of storage (for stack space to support recursion). A marking process that does not require additional stack space was developed by Schorr and Waite (1967). Their method reverses pointers as it traces out linked structures. Then, when the end of a list is reached, the process can follow the pointers back out of the structure.

The most serious problem with the original version of mark-sweep was that it was done too infrequently—only when a program had used all or nearly all of
the heap storage. Mark-sweep in that situation takes a good deal of time, because most of the cells must be traced and marked as being currently used. This causes a significant delay in the progress of the application. Furthermore, the process may yield only a small number of cells that can be placed on the list of available space. This problem has been addressed in a variety of improvements. For example, **incremental mark-sweep** garbage collection occurs more frequently, long before memory is exhausted, making the process more effective in terms of the amount of storage that is reclaimed. Also, the time required for each run of the process is obviously shorter, thus reducing the delay in application execution. Another alternative is to perform the mark-sweep process on parts, rather than all of the memory associated with the application, at different times. This provides the same kinds of improvements as incremental mark-sweep.

Both the marking algorithms for the mark-sweep method and the processes required by the reference counter method can be made more efficient by use of the pointer rotation and slide operations that are described by Suzuki (1982).

**Variable-Size Cells** Managing a heap from which variable-size cells\(^8\) are allocated has all the difficulties of managing one for single-size cells, but also has additional problems. Unfortunately, variable-size cells are required by most programming languages. The additional problems posed by variable-size cell management depend on the method used. If mark-sweep is used, the following additional problems occur:

---

8. The cells have variable sizes because these are abstract cells, which store the values of variables, regardless of their types. Furthermore, a variable could be a structured type.
The initial setting of the indicators of all cells in the heap to indicate that they are garbage is difficult. Because the cells are different sizes, scanning them is a problem. One solution is to require each cell to have the cell size as its first field. Then the scanning can be done, although it takes slightly more space and somewhat more time than its counterpart for fixed-size cells.

The marking process is nontrivial. How can a chain be followed from a pointer if there is no predefined location for the pointer in the pointed-to cell? Cells that do not contain pointers at all are also a problem. Adding an internal pointer to each cell, which is maintained in the background by the run-time system, will work. However, this background maintenance processing adds both space and execution time overhead to the cost of running the program.

Maintaining the list of available space is another source of overhead. The list can begin with a single cell consisting of all available space. Requests for segments simply reduce the size of this block. Reclaimed cells are added to the list. The problem is that before long, the list becomes a long list of various-size segments, or blocks. This slows allocation because requests cause the list to be searched for sufficiently large blocks. Eventually, the list may consist of a large number of very small blocks, which are not large enough for most requests. At this point, adjacent blocks may need to be collapsed into larger blocks. Alternatives to using the first sufficiently large block on the list can shorten the search but require the list to be ordered by block size. In either case, maintaining the list is additional overhead.

If reference counters are used, the first two problems are avoided, but the available-space list-maintenance problem remains.

For a comprehensive study of memory management problems, see Wilson (2005).

For the discussion of type checking, the concept of operands and operators is generalized to include subprograms and assignment statements. Subprograms will be thought of as operators whose operands are their parameters. The assignment symbol will be thought of as a binary operator, with its target variable and its expression being the operands.

Type checking is the activity of ensuring that the operands of an operator are of compatible types. A compatible type is one that either is legal for the operator or is allowed under language rules to be implicitly converted by compiler-generated code (or the interpreter) to a legal type. This automatic conversion is called a coercion. For example, if an int variable and a float variable are added in Java, the value of the int variable is coerced to float and a floating-point add is done.

A type error is the application of an operator to an operand of an inappropriate type. For example, in the original version of C, if an int value was passed
to a function that expected a `float` value, a type error would occur (because compilers for that language did not check the types of parameters).

If all bindings of variables to types are static in a language, then type checking can nearly always be done statically. Dynamic type binding requires type checking at run time, which is called **dynamic type checking**.

Some languages, such as JavaScript and PHP, because of their dynamic type binding, allow only dynamic type checking. It is better to detect errors at compile time than at run time, because the earlier correction is usually less costly. The penalty for static checking is reduced programmer flexibility. Fewer shortcuts and tricks are possible. Such techniques, though, are now generally recognized to be error prone and detrimental to readability.

Type checking is complicated when a language allows a memory cell to store values of different types at different times during execution. Such memory cells can be created with C and C++ unions and the discriminated unions of ML, Haskell, and F#. In these cases, type checking, if done, must be dynamic and requires the run-time system to maintain the type of the current value of such memory cells. So, even though all variables are statically bound to types in languages such as C++, not all type errors can be detected by static type checking.

### 6.13 Strong Typing

One of the ideas in language design that became prominent in the so-called structured-programming revolution of the 1970s was **strong typing**. Strong typing is widely acknowledged as being a highly valuable language characteristic. Unfortunately, it is often loosely defined, and it is often used in computing literature without being defined at all.

A programming language is **strongly typed** if type errors are always detected. This requires that the types of all operands can be determined, either at compile time or at run time. The importance of strong typing lies in its ability to detect all misuses of variables that result in type errors. A strongly typed language also allows the detection, at run time, of uses of the incorrect type values in variables that can store values of more than one type.

C and C++ are not strongly typed languages because both include union types, which are not type checked.

ML is strongly typed, even though the types of some function parameters may not be known at compile time. F# is strongly typed.

Java and C#, although they are based on C++, are nearly strongly typed. Types can be explicitly cast, which could result in a type error. However, there are no implicit ways type errors can go undetected.

The coercion rules of a language have an important effect on the value of type checking. For example, expressions are strongly typed in Java. However, an arithmetic operator with one floating-point operand and one integer operand is legal. The value of the integer operand is coerced to floating-point, and a floating-point operation takes place. This is what is usually intended by the
6.14 Type Equivalence

The idea of type compatibility was defined when the issue of type checking was introduced. The compatibility rules dictate the types of operands that are acceptable for each of the operators and thereby specify the possible type errors of the language. The rules are called compatibility because in some cases the type of an operand can be implicitly converted by the compiler or run-time system to make it acceptable for the operator.

The type compatibility rules are simple and rigid for the predefined scalar types. However, in the cases of structured types, such as arrays and records and some user-defined types, the rules are more complex. Coercion of these types is rare, so the issue is not type compatibility, but type equivalence. That is, two types are equivalent if an operand of one type in an expression is substituted for one of the other type, without coercion. Type equivalence is a strict form of type compatibility—compatibility without coercion. The central issue here is how type equivalence is defined.

The design of the type equivalence rules of a language is important, because it influences the design of the data types and the operations provided for values of those types. With the types discussed here, there are very few predefined operations. Perhaps the most important result of two variables being of equivalent types is that either one can have its value assigned to the other.

There are two approaches to defining type equivalence: name type equivalence and structure type equivalence. Name type equivalence means that two variables have equivalent types if they are defined either in the same declaration or in declarations that use the same type name. Structure type equivalence means that two variables have equivalent types if their types have identical structures. There are some variations of these two approaches, and many languages use combinations of them.

Name type equivalence is easy to implement but is more restrictive. Under a strict interpretation, a variable whose type is a subrange of the integers would...
not be equivalent to an integer type variable. For example, supposing Ada used strict name type equivalence, consider the following Ada code:

```ada
type Indextype is 1..100;
count : Integer;
index : Indextype;
```

The types of the variables `count` and `index` would not be equivalent; `count` could not be assigned to `index` or vice versa.

Another problem with name type equivalence arises when a structured or user-defined type is passed among subprograms through parameters. Such a type must be defined only once, globally. A subprogram cannot state the type of such formal parameters in local terms. This was the case with the original version of Pascal.

Note that to use name type equivalence, all types must have names. Most languages allow users to define types that are anonymous—they do not have names. For a language to use name type equivalence, such types must implicitly be given internal names by the compiler.

Structure type equivalence is more flexible than name type equivalence, but it is more difficult to implement. Under name type equivalence, only the two type names must be compared to determine equivalence. Under structure type equivalence, however, the entire structures of the two types must be compared. This comparison is not always simple. (Consider a data structure that refers to its own type, such as a linked list.) Other questions can also arise. For example, are two record (or `struct`) types equivalent if they have the same structure but different field names? Are two single-dimensional array types in a language that allows lower bounds of array subscript ranges to be set in their declarations equivalent if they have the same element type but have subscript ranges of `0..10` and `1..11`? Are two enumeration types equivalent if they have the same number of components but spell the literals differently?

Another difficulty with structure type equivalence is that it disallows differentiating between types with the same structure. For example, consider the following Ada-like declarations:

```ada
type Celsius = Float;
Fahrenheit = Float;
```

The types of variables of these two types are considered equivalent under structure type equivalence, allowing them to be mixed in expressions, which is surely undesirable in this case, considering the difference indicated by the type’s names. In general, types with different names are likely to be abstractions of different categories of problem values and should not be considered equivalent.

Ada uses a restrictive form of name type equivalence but provides two type constructs, subtypes and derived types, that avoid the problems associated with name type equivalence. A **derived type** is a new type that is based on some previously defined type with which it is not equivalent, although it may have
identical structure. Derived types inherit all the properties of their parent types. Consider the following example:

```ada
type Celsius is new Float;
type Fahrenheit is new Float;
```

The types of variables of these two derived types are not equivalent, although their structures are identical. Furthermore, variables of neither type is type equivalent with any other floating-point type. Literals are exempt from the rule. A literal such as \(3.0\) has the type universal real and is type equivalent to any floating-point type. Derived types can also include range constraints on the parent type, while still inheriting all of the parent’s operations.

An Ada **subtype** is a possibly range-constrained version of an existing type. A subtype is type equivalent with its parent type. For example, consider the following declaration:

```ada
subtype Small_type is Integer range 0..99;
```

The type `Small_type` is equivalent to the type `Integer`.

Note that Ada’s derived types are very different from Ada’s subrange types. For example, consider the following type declarations:

```ada
type Derived_Small_Int is new Integer range 1..100;
subtype Subrange_Small_Int is Integer range 1..100;
```

Variables of both types, `Derived_Small_Int` and `Subrange_Small_Int`, have the same range of legal values and both inherit the operations of `Integer`. However, variables of type `Derived_Small_Int` are not compatible with any `Integer` type. On the other hand, variables of type `Subrange_Small_Int` are compatible with variables and constants of `Integer` type and any subtype of `Integer`.

For variables of an Ada unconstrained array type, structure type equivalence is used. For example, consider the following type declaration and two object declarations:

```ada
type Vector is array (Integer range <>) of Integer;
Vector_1: Vector (1..10);
Vector_2: Vector (11..20);
```

The types of these two objects are equivalent, even though they have different names and different subscript ranges, because for objects of unconstrained array types, structure type equivalence rather than name type equivalence is used. Because both types have 10 elements and the elements of both are of type `Integer`, they are type equivalent.

For constrained anonymous types, Ada uses a highly restrictive form of name type equivalence. Consider the following Ada declarations of constrained anonymous types:

```ada
A : array (1..10) of Integer;
```
In this case, $A$ has an anonymous but unique type assigned by the compiler and unavailable to the program. If we also had

$$B : array \ (1..10) \ of \ Integer;$$

$A$ and $B$ would be of anonymous but distinct and not equivalent types, though they are structurally identical. The multiple declaration

$$C, \ D : array \ (1..10) \ of \ Integer;$$

creates two anonymous types, one for $C$ and one for $D$, which are not equivalent. This declaration is actually treated as if it were the following two declarations:

$$C : array \ (1..10) \ of \ Integer;$$
$$D : array \ (1..10) \ of \ Integer;$$

Note that Ada's form of name type equivalence is more restrictive than the name type equivalence that is defined at the beginning of this section. If we had written instead

$$\text{type List}_10 \ is \ array \ (1..10) \ of \ Integer;$$
$$C, \ D : \text{List}_10;$$

then the types of $C$ and $D$ would be equivalent.

Name type equivalence works well for Ada, in part because all types, except anonymous arrays, are required to have type names (and anonymous types are given internal names by the compiler).

Type equivalence rules for Ada are more rigid than those for languages that have many coercions among types. For example, the two operands of an addition operator in Java can have virtually any combination of numeric types in the language. One of the operands will simply be coerced to the type of the other. But in Ada, there are no coercions of the operands of an arithmetic operator.

C uses both name and structure type equivalence. Every \text{struct}, \text{enum}, and \text{union} declaration creates a new type that is not equivalent to any other type. So, name type equivalence is used for structure, enumeration, and union types. Other non scalar types use structure type equivalence. Array types are equivalent if they have the same type components. Also, if an array type has a constant size, it is equivalent either to other arrays with the same constant size or to with those without a constant size. Note that \text{typedef} in C and C++ does not introduce a new type; it simply defines a new name for an existing type. So, any type defined with \text{typedef} is type equivalent to its parent type. One exception to C using name type equivalence for structures, enumerations, and unions is if two structures, enumerations, or unions are defined in different files, in which case structural type equivalence is used. This is a loophole in the name type equivalence rule to allow equivalence of structures, enumerations, and unions that are defined in different files.
C++ is like C except there is no exception for structures and unions defined in different files.

In languages that do not allow users to define and name types, such as Fortran and COBOL, name equivalence obviously cannot be used.

Object-oriented languages such as Java and C++ bring another kind of type compatibility issue with them. The issue is object compatibility and its relationship to the inheritance hierarchy, which is discussed in Chapter 12.

Type compatibility in expressions is discussed in Chapter 7; type compatibility for subprogram parameters is discussed in Chapter 9.

6.15 Theory and Data Types

Type theory is a broad area of study in mathematics, logic, computer science, and philosophy. It began in mathematics in the early 1900s and later became a standard tool in logic. Any general discussion of type theory is necessarily complex, lengthy, and highly abstract. Even when restricted to computer science, type theory includes such diverse and complex subjects as typed lambda calculus, combinators, the metatheory of bounded quantification, existential types, and higher-order polymorphism. All these topics are far beyond the scope of this book.

In computer science there are two branches of type theory: practical and abstract. The practical branch is concerned with data types in commercial programming languages; the abstract branch primarily focuses on typed lambda calculus, an area of extensive research by theoretical computer scientists over the past half century. This section is restricted to a brief description of some of the mathematical formalisms that underlie data types in programming languages.

A data type defines a set of values and a collection of operations on those values. A type system is a set of types and the rules that govern their use in programs. Obviously, every typed programming language defines a type system. The formal model of a type system of a programming language consists of a set of types and a collection of functions that define the type rules of the language, which are used to determine the type of any expression. A formal system that describes the rules of a type system, attribute grammars, is introduced in Chapter 3.

An alternative model to attribute grammars uses a type map and a collection of functions, not associated with grammar rules, that specify the type rules. A type map is similar to the state of a program used in denotational semantics, consisting of a set of ordered pairs, with the first element of each pair being a variable's name and the second element being its type. A type map is constructed using the type declarations in the program. In a static typed language, the type map need only be maintained during compilation, although it changes as the program is analyzed by the compiler. If any type checking is done dynamically, the type map must be maintained during execution. The concrete version of a type map in a compilation system is the symbol table, constructed primarily
by the lexical and syntax analyzers. Dynamic types sometimes are maintained with tags attached to values or objects.

As stated previously, a data type is a set of values, although in a data type the elements are often ordered. For example, the elements in all enumeration types are ordered. However, in a mathematical set the elements are not ordered. Despite this difference, set operations can be used on data types to describe new data types. The structured data types of programming languages are defined by type operators, or constructors that correspond to set operations. These set operations/type constructors are briefly introduced in the following paragraphs.

A finite mapping is a function from a finite set of values, the domain set, onto values in the range set. Finite mappings model two different categories of types in programming languages, functions and arrays, although in some languages functions are not types. All languages include arrays, which are defined in terms of a mapping function that maps indices to elements in the array. For traditional arrays, the mapping is simple—integer values are mapped to the addresses of array elements; for associative arrays, the mapping is defined by a function that describes a hashing operation. The hashing function maps the keys of the associate arrays, usually character strings,\(^{10}\) to the addresses of the array elements.

A Cartesian, or cross product of \(n\) sets, \(S_1, S_2, \ldots, S_n\), is a set denoted \(S_1 \times S_2 \times \ldots \times S_n\). Each element of the Cartesian product set has one element from each of the constituent sets. So, \(S_1 \times S_2 = \{(x, y) \mid x \text{ is in } S_1 \text{ and } y \text{ is in } S_2\}\). For example, if \(S_1 = \{1, 2\}\) and \(S_2 = \{a, b\}\), \(S_1 \times S_2 = \{(1, a), (1, b), (2, a), (2, b)\}\). A Cartesian product defines tuples in mathematics, which appear in Python, ML, and F# as a data type (see Section 6.5). Cartesian products also model records, or structs, although not exactly. Cartesian products do not have element names, but records require them. For example, consider the following C struct:

```c
struct intFloat {
    int myInt;
    float myFloat;
};
```

This struct defines the Cartesian product type \(\text{int} \times \text{float}\). The names of the elements are \(\text{myInt}\) and \(\text{myFloat}\).

The union of two sets, \(S_1\) and \(S_2\), is defined as \(S_1 \cup S_2 = \{x \mid x \text{ is in } S_1 \text{ or } x \text{ is in } S_2\}\). Set union models the union data types, as described in Section 6.10.

Mathematical subsets are defined by providing a rule that elements must follow. These sets model the subtypes of Ada, although not exactly, because subtypes must consist of contiguous elements of their parent sets. Elements of mathematical sets are unordered, so the model is not perfect.

\(^{10}\) In Ruby and Lua, the associative array keys need not be character strings—they can be any type.
Notice that pointers, defined with type operators, such as `*` in C, are not defined in terms of a set operation.

This concludes our discussion of formalisms in data types, as well as our whole discussion of data types.

**Summary**

The data types of a language are a large part of what determines that language's style and usefulness. Along with control structures, they form the heart of a language.

The primitive data types of most imperative languages include numeric, character, and Boolean types. The numeric types are often directly supported by hardware.

The user-defined enumeration and subrange types are convenient and add to the readability and reliability of programs.

Arrays are part of most programming languages. The relationship between a reference to an array element and the address of that element is given in an access function, which is an implementation of a mapping. Arrays can be either static, as in C++ arrays whose definition includes the `static` specifier; fixed stack-dynamic, as in C functions (without the `static` specifier); fixed heap-dynamic, as with Java's objects; or heap dynamic, as in Perl's arrays. Most languages allow only a few operations on complete arrays.

Records are now included in most languages. Fields of records are specified in a variety of ways. In the case of COBOL, they can be referenced without naming all of the enclosing records, although this is messy to implement and harmful to readability. In several languages that support object-oriented programming, records are supported with objects.

Tuples are similar to records, but do not have names for their constituent parts. They are part of Python, ML, and F#.

Lists are staples of the functional programming languages, but are now also included in Python and C#.

Unions are locations that can store different type values at different times. Discriminated unions include a tag to record the current type value. A free union is one without the tag. Most languages with unions do not have safe designs for them, the exceptions being ML and F#.

Pointers are used for addressing flexibility and to control dynamic storage management. Pointers have some inherent dangers: Dangling pointers are difficult to avoid, and memory leakage can occur.

Reference types, such as those in Java and C#, provide heap management without the dangers of pointers.

Enumeration and record types are relatively easy to implement. Arrays are also straightforward, although array element access is an expensive process when the array has several subscripts. The access function requires one addition and one multiplication for each subscript.
Pointers are relatively easy to implement, if heap management is not considered. Heap management is easy if all cells have the same size but is complicated for variable-size cell allocation and deallocation.

Strong typing is the concept of requiring that all type errors be detected. The value of strong typing is increased reliability.

The type equivalence rules of a language determine what operations are legal among the structured types of a language. Name type equivalence and structure type equivalence are the two fundamental approaches to defining type equivalence.

Type theories have been developed in many areas. In computer science, the practical branch of type theory defines the types and type rules of programming languages. Set theory can be used to model most of the structured data types in programming languages.

**BIBLIOGRAPHIC NOTES**

A wealth of literature exists that is concerned with data type design, use, and implementation. Hoare gives one of the earliest systematic definitions of structured types in Dahl et al. (1972). A general discussion of a wide variety of data types is given in Cleaveland (1986).

Implementing run-time checks on the possible insecurities of Pascal data types is discussed in Fischer and LeBlanc (1980). Most compiler design books, such as Fischer and LeBlanc (1991) and Aho et al. (1986), describe implementation methods for data types, as do the other programming language texts, such as Pratt and Zelkowitz (2001) and Scott (2009). A detailed discussion of the problems of heap management can be found in Tenenbaum et al. (1990). Garbage-collection methods are developed by Schorr and Waite (1967) and Deutsch and Bobrow (1976). A comprehensive discussion of garbage-collection algorithms can be found in Cohen (1981) and Wilson (2005).

**REVIEW QUESTIONS**

1. What is a data type?
2. What are the advantages of the data structure of COBOL over the data structure of Fortran I?
3. What is an abstract data type?
4. Describe the different uses of the type system of a programming language.
5. Define `descriptor` and `object`.
6. What are the two most common structured data types in the imperative languages?
7. What mechanism is used to store negative integers in a computer?
8. What are the four signed integers supported by Java?
9. Define *static*, *fixed stack-dynamic*, *fixed heap-dynamic*, and *heap-dynamic arrays*. What are the advantages of each?

10. What is the disadvantage of ones-complement notation?

11. How are floating-point types represented?

12. What are the different representations for floating-point values supported by Java?

13. What languages support complex data type?

14. What is the disadvantage of a decimal type?

15. Define *precision* and *range*.

16. What is the use of a Boolean type?

17. What are the required entries in a Java array descriptor, and when must they be stored (at compile time or run time)?

18. What is a substring reference?

19. What two classes support strings in Java?

20. Define *limited dynamic length strings* and *dynamic length strings*.

21. What is an immutable object of Java’s `String` class?

22. Define *tuple*.

23. What are the design issues for enumeration types?

24. In what primarily imperative language do lists serve as arrays?

25. What method is used to fetch the internal numeric value of an enumeration variable?

26. What is the difference between C# enumeration types and C++ enumeration types?

27. In what way does Scheme’s `CDR` function modify its parameter?

28. What is the mechanics of a list comprehension in Python?

29. Define *union*, *free union*, and *discriminated union*.

30. Are the unions of F# discriminated?

31. What are the advantages of the enumeration types of F#?

32. What is the design issue for associative arrays?

33. Why are associative arrays called hashes in Perl?

34. What are generic arrays in Java?

35. Why are reference variables in C++ better than pointers for formal parameters?

36. What advantages do Java and C# reference type variables have over the pointers in other languages?

37. Describe the lazy and eager approaches to reclaiming garbage.

38. Why wouldn’t arithmetic on Java and C# references make sense?

39. What is the data type of the subscript of an array in Java?

40. Define *record*.

41. Define strongly typed.
42. Why is Java not strongly typed?
43. What is the syntax of array references?
44. What languages have no type coercions?
45. Why are the unions in C and C++ called free unions?
46. What is name type equivalence?
47. What is a dangling pointer?
48. What is the primary advantage of name type equivalence?
49. What is the primary disadvantage to structure type equivalence?
50. For what types does C use structure type equivalence?
51. What set operation models C’s struct data type?

**Problem Set**

1. What are the arguments for and against representing Boolean values as single bits in memory?
2. How does a decimal value waste memory space?
3. VAX minicomputers use a format for floating-point numbers that is not the same as the IEEE standard. What is this format, and why was it chosen by the designers of the VAX computers? A reference for VAX floating-point representations is Sebesta (1991).
4. Compare the tombstone and lock-and-key methods of avoiding dangling pointers, from the points of view of safety and implementation cost.
5. What disadvantages are there in implicit dereferencing of pointers, but only in certain contexts?
6. Explain all of the differences between Ada’s subtypes and derived types.
7. What significant justification is there for the -> operator in C and C++?
8. What are all of the differences between the enumeration types of C++ and those of Java?
9. Multidimensional arrays can be stored in row major order, as in C++, or in column major order, as in Fortran. Develop the access functions for both of these arrangements for three-dimensional arrays.
10. In the Burroughs Extended ALGOL language, matrices are stored as a single-dimensioned array of pointers to the rows of the matrix, which are treated as single-dimensioned arrays of values. What are the advantages and disadvantages of such a scheme?
11. Analyze and write a comparison of C’s malloc and free functions with C++’s new and delete operators. Use safety as the primary consideration in the comparison.
12. Analyze and write a comparison of using C++ pointers and Java reference variables to refer to fixed heap-dynamic variables. Use safety and convenience as the primary considerations in the comparison.

13. Write a short discussion of what was lost and what was gained in Java’s designers’ decision to not include the pointers of C++.

14. What are the arguments for and against Java’s implicit heap storage recovery, when compared with the explicit heap storage recovery required in C++? Consider real-time systems.

15. What are the arguments for the inclusion of enumeration types in C#, although they were not in the first few versions of Java?

16. What would you expect to be the level of use of pointers in C#? How often will they be used when it is not absolutely necessary?

17. Make two lists of applications of matrices, one for those that require jagged matrices and one for those that require rectangular matrices. Now, argue whether just jagged, just rectangular, or both should be included in a programming language.

18. Compare the string manipulation capabilities of the class libraries of C++, Java, and C#.

19. Look up the definition of strongly typed as given in Gehani (1983) and compare it with the definition given in this chapter. How do they differ?

20. In what way is static type checking better than dynamic type checking?

21. Explain how coercion rules can weaken the beneficial effect of strong typing.

---

**Programming Exercises**

1. Design a set of simple test programs to determine the type compatibility rules of a C compiler to which you have access. Write a report of your findings.

2. Determine whether some C compiler to which you have access implements the `free` function.

3. Write a program that does matrix multiplication in some language that does subscript range checking and for which you can obtain an assembly language or machine language version from the compiler. Determine the number of instructions required for the subscript range checking and compare it with the total number of instructions for the matrix multiplication process.

4. If you have access to a compiler in which the user can specify whether subscript range checking is desired, write a program that does a large number of matrix accesses and time their execution. Run the program with subscript range checking and without it, and compare the times.
5. Write a simple program in C++ to investigate the safety of its enumeration types. Include at least 10 different operations on enumeration types to determine what incorrect or just silly things are legal. Now, write a C# program that does the same things and run it to determine how many of the incorrect or silly things are legal. Compare your results.

6. Write a program in C++ or C# that includes two different enumeration types and has a significant number of operations using the enumeration types. Also write the same program using only integer variables. Compare the readability and predict the reliability differences between the two programs.

7. Write a C program that does a large number of references to elements of two-dimensioned arrays, using only subscripting. Write a second program that does the same operations but uses pointers and pointer arithmetic for the storage-mapping function to do the array references. Compare the time efficiency of the two programs. Which of the two programs is likely to be more reliable? Why?

8. Write a Perl program that uses a hash and a large number of operations on the hash. For example, the hash could store people’s names and their ages. A random-number generator could be used to create three-character names and ages, which could be added to the hash. When a duplicate name was generated, it would cause an access to the hash but not add a new element. Rewrite the same program without using hashes. Compare the execution efficiency of the two. Compare the ease of programming and readability of the two.

9. Write a program in the language of your choice that behaves differently if the language used name equivalence than if it used structural equivalence.

10. For what types of A and B is the simple assignment statement \( A = B \) legal in C++ but not Java?
Expressions and Assignment Statements

7.1 Introduction
7.2 Arithmetic Expressions
7.3 Overloaded Operators
7.4 Type Conversions
7.5 Relational and Boolean Expressions
7.6 Short-Circuit Evaluation
7.7 Assignment Statements
7.8 Mixed-Mode Assignment
Expressions and Assignment Statements

As the title indicates, the topic of this chapter is expressions and assignment statements. The semantics rules that determine the order of evaluation of operators in expressions are discussed first. This is followed by an explanation of a potential problem with operand evaluation order when functions can have side effects. Overloaded operators, both predefined and user defined, are then discussed, along with their effects on the expressions in programs. Next, mixed-mode expressions are described and evaluated. This leads to the definition and evaluation of widening and narrowing type conversions, both implicit and explicit. Relational and Boolean expressions are then discussed, including the process of short-circuit evaluation. Finally, the assignment statement, from its simplest form to all of its variations, is covered, including assignments as expressions and mixed-mode assignments.

Character string pattern-matching expressions were covered as a part of the material on character strings in Chapter 6, so they are not mentioned in this chapter.

7.1 Introduction

Expressions are the fundamental means of specifying computations in a programming language. It is crucial for a programmer to understand both the syntax and semantics of expressions of the language he or she uses. A formal mechanism (BNF) for describing the syntax of expressions was introduced in Chapter 3. In this chapter, the semantics of expressions are discussed.

To understand expression evaluation, it is necessary to be familiar with the orders of operator and operand evaluation. The operator evaluation order of expressions is dictated by the associativity and precedence rules of the language. Although the value of an expression sometimes depends on it, the order of operand evaluation in expressions is often unstated by language designers. This allows implementors to choose the order, which leads to the possibility of programs producing different results in different implementations. Other issues in expression semantics are type mismatches, coercions, and short-circuit evaluation.

The essence of the imperative programming languages is the dominant role of assignment statements. The purpose of these statements is to cause the side effect of changing the values of variables, or the state, of the program. So an integral part of all imperative languages is the concept of variables whose values change during program execution.

Functional languages use variables of a different sort, such as the parameters of functions. These languages also have declaration statements that bind values to names. These declarations are similar to assignment statements, but do not have side effects.

7.2 Arithmetic Expressions

Automatic evaluation of arithmetic expressions similar to those found in mathematics, science, and engineering was one of the primary goals of the first high-level programming languages. Most of the characteristics of arithmetic
expressions in programming languages were inherited from conventions that had evolved in mathematics. In programming languages, arithmetic expressions consist of operators, operands, parentheses, and function calls. An operator can be unary, meaning it has a single operand, binary, meaning it has two operands, or ternary, meaning it has three operands.

In most programming languages, binary operators are infix, which means they appear between their operands. One exception is Perl, which has some operators that are prefix, which means they precede their operands.

The purpose of an arithmetic expression is to specify an arithmetic computation. An implementation of such a computation must cause two actions: fetching the operands, usually from memory, and executing arithmetic operations on those operands. In the following sections, we investigate the common design details of arithmetic expressions.

Following are the primary design issues for arithmetic expressions, all of which are discussed in this section:

• What are the operator precedence rules?
• What are the operator associativity rules?
• What is the order of operand evaluation?
• Are there restrictions on operand evaluation side effects?
• Does the language allow user-defined operator overloading?
• What type mixing is allowed in expressions?

7.2.1 Operator Evaluation Order

The operator precedence and associativity rules of a language dictate the order of evaluation of its operators.

7.2.1.1 Precedence

The value of an expression depends at least in part on the order of evaluation of the operators in the expression. Consider the following expression:

\[ a + b \times c \]

Suppose the variables \( a \), \( b \), and \( c \) have the values 3, 4, and 5, respectively. If evaluated left to right (the addition first and then the multiplication), the result is 35. If evaluated right to left, the result is 23.

Instead of simply evaluating the operators in an expression from left to right or right to left, mathematicians long ago developed the concept of placing operators in a hierarchy of evaluation priorities and basing the evaluation order of expressions partly on this hierarchy. For example, in mathematics, multiplication is considered to be of higher priority than addition, perhaps due to its higher level of complexity. If that convention were applied in the previous example expression, as would be the case in most programming languages, the multiplication would be done first.
The **operator precedence rules** for expression evaluation partially define the order in which the operators of different precedence levels are evaluated. The operator precedence rules for expressions are based on the hierarchy of operator priorities, as seen by the language designer. The operator precedence rules of the common imperative languages are nearly all the same, because they are based on those of mathematics. In these languages, exponentiation has the highest precedence (when it is provided by the language), followed by multiplication and division on the same level, followed by binary addition and subtraction on the same level.

Many languages also include unary versions of addition and subtraction. Unary addition is called the **identity operator** because it usually has no associated operation and thus has no effect on its operand. Ellis and Stroustrup (1990, p. 56), speaking about C++, call it a historical accident and correctly label it useless. Unary minus, of course, changes the sign of its operand. In Java and C#, unary minus also causes the implicit conversion of **short** and **byte** operands to **int** type.

In all of the common imperative languages, the unary minus operator can appear in an expression either at the beginning or anywhere inside the expression, as long as it is parenthesized to prevent it from being next to another operator. For example,

```
a + (- b) * c
```

is legal, but

```
a + - b * c
```

usually is not.

Next, consider the following expressions:

```
- a / b
- a * b
- a ** b
```

In the first two cases, the relative precedence of the unary minus operator and the binary operator is irrelevant—the order of evaluation of the two operators has no effect on the value of the expression. In the last case, however, it does matter.

Of the common programming languages, only Fortran, Ruby, Visual Basic, and Ada have the exponentiation operator. In all four, exponentiation has higher precedence than unary minus, so

```
- A ** B
```

is equivalent to

```
- (A ** B)
```
The precedences of the arithmetic operators of Ruby and the C-based languages are as follows:

<table>
<thead>
<tr>
<th>Highest</th>
<th>Ruby</th>
<th>C-Based Languages</th>
</tr>
</thead>
<tbody>
<tr>
<td>**</td>
<td>**</td>
<td>postfix ++, --</td>
</tr>
<tr>
<td>unary +, -</td>
<td>unary +, -</td>
<td>prefix ++, --, unary +, -</td>
</tr>
<tr>
<td>*, /, %</td>
<td>*, /, %</td>
<td>binary +, -</td>
</tr>
</tbody>
</table>

| Lowest      | binary +, - | binary +, - |

The ** operator is exponentiation. The % operator takes two integer operands and yields the remainder of the first after division by the second.\(^1\) The ++ and -- operators of the C-based languages are described in Section 7.7.4.

APL is odd among languages because it has a single level of precedence, as illustrated in the next section.

Precedence accounts for only some of the rules for the order of operator evaluation; associativity rules also affect it.

### 7.2.1.2 Associativity

Consider the following expression:

\[
a - b + c - d
\]

If the addition and subtraction operators have the same level of precedence, as they do in programming languages, the precedence rules say nothing about the order of evaluation of the operators in this expression.

When an expression contains two adjacent\(^2\) occurrences of operators with the same level of precedence, the question of which operator is evaluated first is answered by the associativity rules of the language. An operator can have either left or right associativity, meaning that when there are two adjacent operators with the same precedence, the left operator is evaluated first or the right operator is evaluated first, respectively.

Associativity in common languages is left to right, except that the exponentiation operator (when provided) sometimes associates right to left. In the Java expression

\[
a - b + c
\]

the left operator is evaluated first.

---

1. In versions of C before C99, the % operator was implementation dependent in some situations, because division was also implementation dependent.

2. We call operators “adjacent” if they are separated by a single operand.
Exponentiation in Fortran and Ruby is right associative, so in the expression

\[ A \text{ ** } B \text{ ** } C \]

the right operator is evaluated first.

In Visual Basic, the exponentiation operator, \(^\), is left associative.

The associativity rules for a few common languages are given here:

<table>
<thead>
<tr>
<th>Language</th>
<th>Associativity Rule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ruby</td>
<td>Left: *, /, +, -</td>
</tr>
<tr>
<td></td>
<td>Right: **</td>
</tr>
<tr>
<td>C-based languages</td>
<td>Left: *, /, %, binary +, binary -</td>
</tr>
<tr>
<td></td>
<td>Right: ++, --, unary -, unary +</td>
</tr>
</tbody>
</table>

As stated in Section 7.2.1.1, in APL, all operators have the same level of precedence. Thus, the order of evaluation of operators in APL expressions is determined entirely by the associativity rule, which is right to left for all operators. For example, in the expression

\[ A \times B + C \]

the addition operator is evaluated first, followed by the multiplication operator \((\times\) is the APL multiplication operator). If \(A\) were 3, \(B\) were 4, and \(C\) were 5, then the value of this APL expression would be 27.

Many compilers for the common languages make use of the fact that some arithmetic operators are mathematically associative, meaning that the associativity rules have no impact on the value of an expression containing only those operators. For example, addition is mathematically associative, so in mathematics the value of the expression

\[ A + B + C \]

does not depend on the order of operator evaluation. If floating-point operations for mathematically associative operations were also associative, the compiler could use this fact to perform some simple optimizations. Specifically, if the compiler is allowed to reorder the evaluation of operators, it may be able to produce slightly faster code for expression evaluation. Compilers commonly do these kinds of optimizations.

Unfortunately, in a computer, both floating-point representations and floating-point arithmetic operations are only approximations of their mathematical counterparts (because of size limitations). The fact that a mathematical operator is associative does not necessarily imply that the corresponding floating-point operation is associative. In fact, only if all the operands and intermediate results can be exactly represented in floating-point notation will the process be precisely associative. For example, there are pathological situations
7.2 Arithmetic Expressions

in which integer addition on a computer is not associative. For example, suppose that a program must evaluate the expression

\[ A + B + C + D \]

and that \( A \) and \( C \) are very large positive numbers, and \( B \) and \( D \) are negative numbers with very large absolute values. In this situation, adding \( B \) to \( A \) does not cause an overflow exception, but adding \( C \) to \( A \) does. Likewise, adding \( C \) to \( B \) does not cause overflow, but adding \( D \) to \( B \) does. Because of the limitations of computer arithmetic, addition is catastrophically nonassociative in this case. Therefore, if the compiler reorders these addition operations, it affects the value of the expression. This problem, of course, can be avoided by the programmer, assuming the approximate values of the variables are known. The programmer can specify the expression in two parts (in two assignment statements), ensuring that overflow is avoided. However, this situation can arise in far more subtle ways, in which the programmer is less likely to notice the order dependence.

7.2.1.3 Parentheses

Programmers can alter the precedence and associativity rules by placing parentheses in expressions. A parenthesized part of an expression has precedence over its adjacent unparenthesized parts. For example, although multiplication has precedence over addition, in the expression

\[(A + B) \times C\]

the addition will be evaluated first. Mathematically, this is perfectly natural. In this expression, the first operand of the multiplication operator is not available until the addition in the parenthesized subexpression is evaluated. Also, the expression from Section 7.2.1.2 could be specified as

\[(A + B) + (C + D)\]

to avoid overflow.

Languages that allow parentheses in arithmetic expressions could dispense with all precedence rules and simply associate all operators left to right or right to left. The programmer would specify the desired order of evaluation with parentheses. This approach would be simple because neither the author nor the readers of programs would need to remember any precedence or associativity rules. The disadvantage of this scheme is that it makes writing expressions more tedious, and it also seriously compromises the readability of the code. Yet this was the choice made by Ken Iverson, the designer of APL.

7.2.1.4 Ruby Expressions

Recall that Ruby is a pure object-oriented language, which means, among other things, that every data value, including literals, is an object. Ruby supports the collection of arithmetic and logic operations that are included in the C-based
languages. What sets Ruby apart from the C-based languages in the area of expressions is that all of the arithmetic, relational, and assignment operators, as well as array indexing, shifts, and bitwise logic operators, are implemented as methods. For example, the expression $a + b$ is a call to the $+$ method of the object referenced by $a$, passing the object referenced by $b$ as a parameter.

One interesting result of the implementation of operators as methods is that they can be overridden by application programs. Therefore, these operators can be redefined. While it is often not useful to redefine operators for predefined types, it is useful, as we will see in Section 7.3, to define predefined operators for user-defined types, which can be done with operator overloading in some languages.

7.2.1.5 Expressions in Lisp

As is the case with Ruby, all arithmetic and logic operations in Lisp are performed by subprograms. But in Lisp, the subprograms must be explicitly called. For example, to specify the C expression $a + b \times c$ in Lisp, one must write the following expression:

$$+ a (* b c)$$

In this expression, + and * are the names of functions.

7.2.1.6 Conditional Expressions

if-then-else statements can be used to perform a conditional expression assignment. For example, consider

```c
if (count == 0)
  average = 0;
else
  average = sum / count;
```

In the C-based languages, this code can be specified more conveniently in an assignment statement using a conditional expression, which has the following form:

```
expression_1 ? expression_2 : expression_3
```

where expression_1 is interpreted as a Boolean expression. If expression_1 evaluates to true, the value of the whole expression is the value of expression_2; otherwise, it is the value of expression_3. For example, the effect of the example if-then-else can be achieved with the following assignment statement, using a conditional expression:

```c
average = (count == 0) ? 0 : sum / count;
```

3. When a list is interpreted as code in Lisp, the first element is the function name and others are parameters to the function.
In effect, the question mark denotes the beginning of the **then** clause, and the colon marks the beginning of the **else** clause. Both clauses are mandatory. Note that ? is used in conditional expressions as a ternary operator.

Conditional expressions can be used anywhere in a program (in a C-based language) where any other expression can be used. In addition to the C-based languages, conditional expressions are provided in Perl, JavaScript, and Ruby.

### 7.2.2 Operand Evaluation Order

A less commonly discussed design characteristic of expressions is the order of evaluation of operands. Variables in expressions are evaluated by fetching their values from memory. Constants are sometimes evaluated the same way. In other cases, a constant may be part of the machine language instruction and not require a memory fetch. If an operand is a parenthesized expression, all of the operators it contains must be evaluated before its value can be used as an operand.

If neither of the operands of an operator has side effects, then operand evaluation order is irrelevant. Therefore, the only interesting case arises when the evaluation of an operand does have side effects.

#### 7.2.2.1 Side Effects

A **side effect** of a function, naturally called a functional side effect, occurs when the function changes either one of its parameters or a global variable. (A global variable is declared outside the function but is accessible in the function.)

Consider the following expression:

\[ a + \text{fun}(a) \]

If fun does not have the side effect of changing \( a \), then the order of evaluation of the two operands, \( a \) and \( \text{fun}(a) \), has no effect on the value of the expression. However, if \( \text{fun} \) changes \( a \), there is an effect. Consider the following situation: \( \text{fun} \) returns 10 and changes the value of its parameter to 20. Suppose we have the following:

\[
\begin{align*}
a &= 10; \\
b &= a + \text{fun}(a); \\
\end{align*}
\]

Then, if the value of \( a \) is fetched first (in the expression evaluation process), its value is 10 and the value of the expression is 20. But if the second operand is evaluated first, then the value of the first operand is 20 and the value of the expression is 30.

The following C program illustrates the same problem when a function changes a global variable that appears in an expression:

```c
int a = 5; 
int fun1() { 
```
a = 17;
return 3;
} /* end of fun1 */

void main() {
    a = a + fun1();
} /* end of main */

The value computed for a in main depends on the order of evaluation of the operands in the expression a + fun1(). The value of a will be either 8 (if a is evaluated first) or 20 (if the function call is evaluated first).

Note that functions in mathematics do not have side effects, because there is no notion of variables in mathematics. The same is true for functional programming languages. In both mathematics and functional programming languages, functions are much easier to reason about and understand than those in imperative languages, because their context is irrelevant to their meaning.

There are two possible solutions to the problem of operand evaluation order and side effects. First, the language designer could disallow function evaluation from affecting the value of expressions by simply disallowing functional side effects. Second, the language definition could state that operands in expressions are to be evaluated in a particular order and demand that implementors guarantee that order.

Disallowing functional side effects in the imperative languages is difficult, and it eliminates some flexibility for the programmer. Consider the case of C and C++, which have only functions, meaning that all subprograms return one value. To eliminate the side effects of two-way parameters and still provide subprograms that return more than one value, the values would need to be placed in a struct and the struct returned. Access to globals in functions would also have to be disallowed. However, when efficiency is important, using access to global variables to avoid parameter passing is an important method of increasing execution speed. In compilers, for example, global access to data such as the symbol table is commonplace.

The problem with having a strict evaluation order is that some code optimization techniques used by compilers involve reordering operand evaluations. A guaranteed order disallows those optimization methods when function calls are involved. There is, therefore, no perfect solution, as is borne out by actual language designs.

The Java language definition guarantees that operands appear to be evaluated in left-to-right order, eliminating the problem discussed in this section.

### 7.2.2.2 Referential Transparency and Side Effects

The concept of referential transparency is related to and affected by functional side effects. A program has the property of referential transparency if any two expressions in the program that have the same value can be substituted for one another anywhere in the program, without affecting the action of the program. The value of a referentially transparent function depends entirely on
The connection of referential transparency and functional side effects is illustrated by the following example:

```plaintext
result1 = (fun(a) + b) / (fun(a) - c);
temp = fun(a);
result2 = (temp + b) / (temp - c);
```

If the function `fun` has no side effects, `result1` and `result2` will be equal, because the expressions assigned to them are equivalent. However, suppose `fun` has the side effect of adding 1 to either `b` or `c`. Then `result1` would not be equal to `result2`. So, that side effect violates the referential transparency of the program in which the code appears.

There are several advantages to referentially transparent programs. The most important of these is that the semantics of such programs is much easier to understand than the semantics of programs that are not referentially transparent. Being referentially transparent makes a function equivalent to a mathematical function, in terms of ease of understanding.

Because they do not have variables, programs written in pure functional languages are referentially transparent. Functions in a pure functional language cannot have state, which would be stored in local variables. If such a function uses a value from outside the function, that value must be a constant, since there are no variables. Therefore, the value of the function depends on the values of its parameters.

Referential transparency will be further discussed in Chapter 15.

### 7.3 Overloaded Operators

Arithmetic operators are often used for more than one purpose. For example, `+` usually is used to specify integer addition and floating-point addition. Some languages—Java, for example—also use it for string concatenation. This multiple use of an operator is called **operator overloading** and is generally thought to be acceptable, as long as neither readability nor reliability suffers.

As an example of the possible dangers of overloading, consider the use of the ampersand (`&`) in C++. As a binary operator, it specifies a bitwise logical AND operation. As a unary operator, however, its meaning is totally different. As a unary operator with a variable as its operand, the expression value is the address of that variable. In this case, the ampersand is called the address-of operator. For example, the execution of

```plaintext
x = &y;
```

Furthermore, the value of the function cannot depend on the order in which its parameters are evaluated.
causes the address of y to be placed in x. There are two problems with this multiple use of the ampersand. First, using the same symbol for two completely unrelated operations is detrimental to readability. Second, the simple keying error of leaving out the first operand for a bitwise AND operation can go undetected by the compiler, because it is interpreted as an address-of operator. Such an error may be difficult to diagnose.

Virtually all programming languages have a less serious but similar problem, which is often due to the overloading of the minus operator. The problem is only that the compiler cannot tell if the operator is meant to be binary or unary.\(^5\) So once again, failure to include the first operand when the operator is meant to be binary cannot be detected as an error by the compiler. However, the meanings of the two operations, unary and binary, are at least closely related, so readability is not adversely affected.

Some languages that support abstract data types (see Chapter 11), for example, C++, C#, and F#, allow the programmer to further overload operator symbols. For instance, suppose a user wants to define the * operator between a scalar integer and an integer array to mean that each element of the array is to be multiplied by the scalar. Such an operator could be defined by writing a function subprogram named * that performs this new operation. The compiler will choose the correct meaning when an overloaded operator is specified, based on the types of the operands, as with language-defined overloaded operators. For example, if this new definition for * is defined in a C# program, a C# compiler will use the new definition for * whenever the * operator appears with a simple integer as the left operand and an integer array as the right operand.

When sensibly used, user-defined operator overloading can aid readability. For example, if + and * are overloaded for a matrix abstract data type and A, B, C, and D are variables of that type, then

\[
A * B + C * D
\]

can be used instead of

\[
\text{MatrixAdd} (\text{MatrixMult}(A, B), \text{MatrixMult}(C, D))
\]

On the other hand, user-defined overloading can be harmful to readability. For one thing, nothing prevents a user from defining * to mean multiplication. Furthermore, seeing an * operator in a program, the reader must find both the types of the operands and the definition of the operator to determine its meaning. Any or all of these definitions could be in other files.

Another consideration is the process of building a software system from modules created by different groups. If the different groups overloaded the same operators in different ways, these differences would obviously need to be eliminated before putting the system together.

---

5. ML alleviates this problem by using different symbols for unary and binary minus operators, tilde (~) for unary and dash (−) for binary.
C++ has a few operators that cannot be overloaded. Among these are the class or structure member operator (.) and the scope resolution operator (::). Interestingly, operator overloading was one of the C++ features that was not copied into Java. However, it did reappear in C#.

The implementation of user-defined operator overloading is discussed in Chapter 9.

### 7.4 Type Conversions

Type conversions are either narrowing or widening. A narrowing conversion converts a value to a type that cannot store even approximations of all of the values of the original type. For example, converting a `double` to a `float` in Java is a narrowing conversion, because the range of `double` is much larger than that of `float`. A widening conversion converts a value to a type that can include at least approximations of all of the values of the original type. For example, converting an `int` to a `float` in Java is a widening conversion. Widening conversions are nearly always safe, meaning that the approximate magnitude of the converted value is maintained. Narrowing conversions are not always safe—sometimes the magnitude of the converted value is changed in the process. For example, if the floating-point value 1.3E25 is converted to an integer in a Java program, the result will not be in any way related to the original value.

Although widening conversions are usually safe, they can result in reduced accuracy. In many language implementations, although integer-to-floating-point conversions are widening conversions, some precision may be lost. For example, in many cases, integers are stored in 32 bits, which allows at least 9 decimal digits of precision. But floating-point values are also stored in 32 bits, with only about seven decimal digits of precision (because of the space used for the exponent). So, integer-to-floating-point widening can result in the loss of two digits of precision.

Coercions of nonprimitive types are, of course, more complex. In Chapter 5, the complications of assignment compatibility of array and record types were discussed. There is also the question of what parameter types and return types of a method allow it to override a method in a superclass—only when the types are the same, or also some other situations. That issue, as well as the concept of subclasses as subtypes, are discussed in Chapter 12.

Type conversions can be either explicit or implicit. The following two subsections discuss these kinds of type conversions.

#### 7.4.1 Coercion in Expressions

One of the design decisions concerning arithmetic expressions is whether an operator can have operands of different types. Languages that allow such expressions, which are called mixed-mode expressions, must define conventions for implicit operand type conversions because computers do not have binary operations that take operands of different types. Recall that in Chapter 5,
coercion was defined as an implicit type conversion that is initiated by the compiler or runtime system. Type conversions explicitly requested by the programmer are referred to as explicit conversions, or casts, not coercions.

Although some operator symbols may be overloaded, we assume that a computer system, either in hardware or in some level of software simulation, has an operation for each operand type and operator defined in the language. For overloaded operators in a language that uses static type binding, the compiler chooses the correct type of operation on the basis of the types of the operands. When the two operands of an operator are not of the same type and that is legal in the language, the compiler must choose one of them to be coerced and generate the code for that coercion. In the following discussion, the coercion design choices of several common languages are examined.

Language designers are not in agreement on the issue of coercions in arithmetic expressions. Those against a broad range of coercions are concerned with the reliability problems that can result from such coercions, because they reduce the benefits of type checking. Those who would rather include a wide range of coercions are more concerned with the loss in flexibility that results from restrictions. The issue is whether programmers should need to be concerned with this category of errors or whether the compiler should detect them.

As a simple illustration of the problem, consider the following Java code:

```java
int a;
float b, c, d;
...
d = b * a;
```

Assume that the second operand of the multiplication operator was supposed to be \( c \), but because of a keying error it was typed as \( a \). Because mixed-mode expressions are legal in Java, the compiler would not detect this as an error. It would simply insert code to coerce the value of the \texttt{int} operand, \( a \), to \texttt{float}. If mixed-mode expressions were not legal in Java, this keying error would have been detected by the compiler as a type error.

Because error detection is reduced when mixed-mode expressions are allowed, F# and ML do not allow them. For example, they do not allow mixing of integer and floating-point operands in expressions.

In most of the other common languages, there are no restrictions on mixed-mode arithmetic expressions.

The C-based languages have integer types that are smaller than the \texttt{int} type. In Java, these are \texttt{byte} and \texttt{short}. Operands of all of these types are coerced to \texttt{int} whenever virtually any operator is applied to them. So, while data can be stored in variables of these types, it cannot be manipulated before conversion to a larger type. For example, consider the following Java code:

```java
int a;
float b, c, d;
...
d = b * a;
```

Assume that the second operand of the multiplication operator was supposed to be \( c \), but because of a keying error it was typed as \( a \). Because mixed-mode expressions are legal in Java, the compiler would not detect this as an error. It would simply insert code to coerce the value of the \texttt{int} operand, \( a \), to \texttt{float}. If mixed-mode expressions were not legal in Java, this keying error would have been detected by the compiler as a type error.

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```java
int a;
float b, c, d;
...
d = b * a;
```
7.4 Type Conversions

```java
byte a, b, c;
...
a = b + c;
```

The values of `b` and `c` are coerced to `int` and an `int` addition is performed. Then, the sum is converted to `byte` and put in `a`. Given the large size of the memories of contemporary computers, there is little incentive to use `byte` and `short`, unless a large number of them must be stored.

### 7.4.2 Explicit Type Conversion

Most languages provide some capability for doing explicit conversions, both widening and narrowing. In some cases, warning messages are produced when an explicit narrowing conversion results in a significant change to the value of the object being converted.

In the C-based languages, explicit type conversions are called casts. To specify a cast, the desired type is placed in parentheses just before the expression to be converted, as in

```
(int) angle
```

One of the reasons for the parentheses around the type name in these conversions is that the first of these languages, C, has several two-word type names, such as `long int`.

In ML and F#, the casts have the syntax of function calls. For example, in F# we could have the following:

```fsharp
float(sum)
```

### 7.4.3 Errors in Expressions

A number of errors can occur during expression evaluation. If the language requires type checking, either static or dynamic, then operand type errors cannot occur. The errors that can occur because of coercions of operands in expressions have already been discussed. The other kinds of errors are due to the limitations of computer arithmetic and the inherent limitations of arithmetic. The most common error occurs when the result of an operation cannot be represented in the memory cell where it must be stored. This is called overflow or underflow, depending on whether the result was too large or too small. One limitation of arithmetic is that division by zero is disallowed. Of course, the fact that it is not mathematically allowed does not prevent a program from attempting to do it.

Floating-point overflow, underflow, and division by zero are examples of run-time errors, which are sometimes called exceptions. Language facilities that allow programs to detect and deal with exceptions are discussed in Chapter 14.
Chapter 7  Expressions and Assignment Statements

7.5 Relational and Boolean Expressions

In addition to arithmetic expressions, programming languages support relational and Boolean expressions.

7.5.1 Relational Expressions

A relational operator is an operator that compares the values of its two operands. A relational expression has two operands and one relational operator. The value of a relational expression is Boolean, except when Boolean is not a type included in the language. The relational operators are often overloaded for a variety of types. The operation that determines the truth or falsehood of a relational expression depends on the operand types. It can be simple, as for integer operands, or complex, as for character string operands. Typically, the types of the operands that can be used for relational operators are numeric types, strings, and enumeration types.

The syntax of the relational operators for equality and inequality differs among some programming languages. For example, for inequality, the C-based languages use `!=`, Lua uses `~=` or `<>`, and Fortran 95+ uses `.NE.` or `<>`, and ML and F# use `<>

JavaScript and PHP have two additional relational operators, `===` and `!==`. These are similar to their relatives, `==` and `!=`, but prevent their operands from being coerced. For example, the expression

"7" == 7

is true in JavaScript, because when a string and a number are the operands of a relational operator, the string is coerced to a number. However,

"7" === 7

is false, because no coercion is done on the operands of this operator.

Ruby uses `==` for the equality relational operator that uses coercions, and `eql?` for equality with no coercions. Ruby uses `===` only in the `when` clause of its `case` statement, as discussed in Chapter 8.

The relational operators always have lower precedence than the arithmetic operators, so that in expressions such as

a + 1 > 2 * b

the arithmetic expressions are evaluated first.

7.5.2 Boolean Expressions

Boolean expressions consist of Boolean variables, Boolean constants, relational expressions, and Boolean operators. The operators usually include those for the AND, OR, and NOT operations, and sometimes for exclusive OR and
equivalence. Boolean operators usually take only Boolean operands (Boolean variables, Boolean literals, or relational expressions) and produce Boolean values.

In the mathematics of Boolean algebras, the OR and AND operators must have equal precedence. However, the C-based languages assign a higher precedence to AND than OR. Perhaps this resulted from the baseless correlation of multiplication with AND and of addition with OR, which would naturally assign higher precedence to AND.

Because arithmetic expressions can be the operands of relational expressions, and relational expressions can be the operands of Boolean expressions, the three categories of operators must be placed in different precedence levels, relative to each other.

The precedence of the arithmetic, relational, and Boolean operators in the C-based languages is as follows:

| Highest | postfix ++, -- |
|         | unary +, unary -, prefix ++, --, ! |
|         | *, /, % |
|         | binary +, binary - |
|         | <, >, <=, >= |
|         | =, != |
|         | && |
| Lowest  | || |

Versions of C prior to C99 are odd among the popular imperative languages in that they have no Boolean type and thus no Boolean values. Instead, numeric values are used to represent Boolean values. In place of Boolean operands, scalar variables (numeric or character) and constants are used, with zero considered false and all nonzero values considered true. The result of evaluating such an expression is an integer, with the value 0 if false and 1 if true. Arithmetic expressions can also be used for Boolean expressions in C99 and C++.

One odd result of C’s design of relational expressions is that the following expression is legal:

```c
a > b > c
```

The leftmost relational operator is evaluated first because the relational operators of C are left associative, producing either 0 or 1. Then, this result is compared with the variable c. There is never a comparison between b and c in this expression.

Some languages, including Perl and Ruby, provide two sets of the binary logic operators, && and **and** for AND and || and **or** for OR. One difference between && and **and** (and || and **or**) is that the spelled versions have lower precedence. Also, **and** and **or** have equal precedence, but && has higher precedence than ||.
When the nonarithmetic operators of the C-based languages are included, there are more than 40 operators and at least 14 different levels of precedence. This is clear evidence of the richness of the collections of operators and the complexity of expressions possible in these languages.

Readability dictates that a language should include a Boolean type, as was stated in Chapter 6, rather than simply using numeric types in Boolean expressions. Some error detection is lost in the use of numeric types for Boolean operands, because any numeric expression, whether intended or not, is a legal operand to a Boolean operator. In the other imperative languages, any non-Boolean expression used as an operand of a Boolean operator is detected as an error.

7.6 Short-Circuit Evaluation

A short-circuit evaluation of an expression is one in which the result is determined without evaluating all of the operands and/or operators. For example, the value of the arithmetic expression

\[(13 * a) * (b / 13 - 1)\]

is independent of the value of \((b / 13 - 1)\) if \(a\) is 0, because \(0 * x = 0\) for any \(x\). So, when \(a\) is 0, there is no need to evaluate \((b / 13 - 1)\) or perform the second multiplication. However, in arithmetic expressions, this shortcut is not easily detected during execution, so it is never taken.

The value of the Boolean expression

\[(a >= 0) && (b < 10)\]

is independent of the second relational expression if \(a\) is less than zero, because the expression \(\text{FALSE} && (b < 10)\) is \text{FALSE} for all values of \(b\). So, when \(a\) is less than zero, there is no need to evaluate \(b\), the constant 10, the second relational expression, or the && operation. Unlike the case of arithmetic expressions, this shortcut can be easily discovered during execution.

To illustrate a potential problem with non-short-circuit evaluation of Boolean expressions, suppose Java did not use short-circuit evaluation. A table lookup loop could be written using the \textbf{while} statement. One simple version of Java code for such a lookup, assuming that \texttt{list}, which has \texttt{listlen} elements, is the array to be searched and \texttt{key} is the searched-for value, is

```java
index = 0;
while ((index < listlen) && (list[index] != key))
    index = index + 1;
```

If evaluation is not short-circuit, both relational expressions in the Boolean expression of the \textbf{while} statement are evaluated, regardless of the value of the
first. Thus, if key is not in list, the program will terminate with a subscript out-of-range exception. The same iteration that has index == listlen will reference list[listlen], which causes the indexing error because list is declared to have listlen-1 as an upper-bound subscript value.

If a language provides short-circuit evaluation of Boolean expressions and it is used, this is not a problem. In the preceding example, a short-circuit evaluation scheme would evaluate the first operand of the AND operator, but it would skip the second operand if the first operand is false.

A language that provides short-circuit evaluations of Boolean expressions and also has side effects in expressions allows subtle errors to occur. Suppose that short-circuit evaluation is used on an expression and part of the expression that contains a side effect is not evaluated; then the side effect will occur only in complete evaluations of the whole expression. If program correctness depends on the side effect, short-circuit evaluation can result in a serious error. For example, consider the Java expression

\[(a > b) \text{ || } ((b++) / 3)\]

In this expression, b is changed (in the second arithmetic expression) only when \(a \leq b\). If the programmer assumed \(b\) would be changed every time this expression is evaluated during execution (and the program’s correctness depends on it), the program will fail.

In the C-based languages, the usual AND and OR operators, && and ||, respectively, are short-circuit. However, these languages also have bitwise AND and OR operators, & and |, respectively, that can be used on Boolean-valued operands and are not short-circuit. Of course, the bitwise operators are only equivalent to the usual Boolean operators if all operands are restricted to being either 0 (for false) or 1 (for true).

All of the logical operators of Ruby, Perl, ML, F#, and Python are short-circuit evaluated.

### 7.7 Assignment Statements

As we have previously stated, the assignment statement is one of the central constructs in imperative languages. It provides the mechanism by which the user can dynamically change the bindings of values to variables. In the following section, the simplest form of assignment is discussed. Subsequent sections describe a variety of alternatives.

#### 7.7.1 Simple Assignments

Nearly all programming languages currently being used use the equal sign for the assignment operator. All of these must use something different from an equal sign for the equality relational operator to avoid confusion with their assignment operator.
ALGOL 60 pioneered the use of := as the assignment operator, which avoids the confusion of assignment with equality. Ada also uses this assignment operator.

The design choices of how assignments are used in a language have varied widely. In some languages, such as Fortran and Ada, an assignment can appear only as a stand-alone statement, and the destination is restricted to a single variable. There are, however, many alternatives.

### 7.7.2 Conditional Targets

Perl allows conditional targets on assignment statements. For example, consider

```
($flag ? $count1 : $count2) = 0;
```

which is equivalent to

```
if ($flag) {
    $count1 = 0;
} else {
    $count2 = 0;
}
```

### 7.7.3 Compound Assignment Operators

A **compound assignment operator** is a shorthand method of specifying a commonly needed form of assignment. The form of assignment that can be abbreviated with this technique has the destination variable also appearing as the first operand in the expression on the right side, as in

```
a = a + b
```

Compound assignment operators were introduced by ALGOL 68, were later adopted in a slightly different form by C, and are part of the other C-based languages, as well as Perl, JavaScript, Python, and Ruby. The syntax of these assignment operators is the catenation of the desired binary operator to the = operator. For example,

```
sum += value;
```

is equivalent to

```
sum = sum + value;
```

The languages that support compound assignment operators have versions for most of their binary operators.
7.7.4 Unary Assignment Operators

The C-based languages, Perl, and JavaScript include two special unary arithmetic operators that are actually abbreviated assignments. They combine increment and decrement operations with assignment. The operators `++` for increment and `--` for decrement can be used either in expressions or to form stand-alone single-operator assignment statements. They can appear either as prefix operators, meaning that they precede the operands, or as postfix operators, meaning that they follow the operands. In the assignment statement

```c
sum = ++ count;
```

the value of `count` is incremented by 1 and then assigned to `sum`. This operation could also be stated as

```c
count = count + 1;
sum = count;
```

If the same operator is used as a postfix operator, as in

```c
sum = count ++;
```

the assignment of the value of `count` to `sum` occurs first; then `count` is incremented. The effect is the same as that of the two statements

```c
sum = count;
count = count + 1;
```

An example of the use of the unary increment operator to form a complete assignment statement is

```c
count ++;
```

which simply increments `count`. It does not look like an assignment, but it certainly is one. It is equivalent to the statement

```c
count = count + 1;
```

When two unary operators apply to the same operand, the association is right to left. For example, in

```c
- count ++
count
```

is first incremented and then negated. So, it is equivalent to

```c
- (count ++)
```

rather than

```c
(- count) ++
```
Assignment as an Expression

In the C-based languages, Perl, and JavaScript, the assignment statement produces a result, which is the same as the value assigned to the target. It can therefore be used as an expression and as an operand in other expressions. This design treats the assignment operator much like any other binary operator, except that it has the side effect of changing its left operand. For example, in C, it is common to write statements such as

```c
while ((ch = getchar()) != EOF) {...}
```

In this statement, the next character from the standard input file, usually the keyboard, is gotten with `getchar` and assigned to the variable `ch`. The result, or value assigned, is then compared with the constant `EOF`. If `ch` is not equal to `EOF`, the compound statement `{...}` is executed. Note that the assignment must be parenthesized—in the languages that support assignment as an expression, the precedence of the assignment operator is lower than that of the relational operators. Without the parentheses, the new character would be compared with `EOF` first. Then, the result of that comparison, either 0 or 1, would be assigned to `ch`.

The disadvantage of allowing assignment statements to be operands in expressions is that it provides yet another kind of expression side effect. This type of side effect can lead to expressions that are difficult to read and understand. An expression with any kind of side effect has this disadvantage. Such an expression cannot be read as an expression, which in mathematics is a denotation of a value, but only as a list of instructions with an odd order of execution. For example, the expression

```c
a = b + (c = d / b) - 1
```

denotes the instructions

- Assign `d / b` to `c`
- Assign `b + c` to `temp`
- Assign `temp - 1` to `a`

Note that the treatment of the assignment operator as any other binary operator allows the effect of multiple-target assignments, such as

```c
sum = count = 0;
```

in which `count` is first assigned the zero, and then `count`'s value is assigned to `sum`. This form of multiple-target assignments is also legal in Python.

There is a loss of error detection in the C design of the assignment operation that frequently leads to program errors. In particular, if we type

```c
if  (x = y) ...  
```

instead of

```c
if  (x == y) ...  
```
which is an easily made mistake, it is not detectable as an error by the compiler. Rather than testing a relational expression, the value that is assigned to \( x \) is tested (in this case, it is the value of \( y \) that reaches this statement). This is actually a result of two design decisions: allowing assignment to behave like an ordinary binary operator and using two very similar operators, \( = \) and \( == \), to have completely different meanings. This is another example of the safety deficiencies of C and C++ programs. Note that Java and C# allow only boolean expressions in their if statements, disallowing this problem.

### 7.7.6 Multiple Assignments

Several recent programming languages, including Perl, Ruby, and Lua, provide multiple-target, multiple-source assignment statements. For example, in Perl one can write

\[
($first, \, $second, \, $third) = (20, \, 40, \, 60);
\]

The semantics is that 20 is assigned to $first, 40 is assigned to $second, and 60 is assigned to $third. If the values of two variables must be interchanged, this can be done with a single assignment, as with

\[
($first, \, $second) = ($second, \, $first);
\]

This correctly interchanges the values of $first and $second, without the use of a temporary variable (at least one created and managed by the programmer).

The syntax of the simplest form of Ruby’s multiple assignment is similar to that of Perl, except the left and right sides are not parenthesized. Also, Ruby includes a few more elaborate versions of multiple assignments, which are not discussed here.

### 7.7.7 Assignment in Functional Programming Languages

All of the identifiers used in pure functional languages and some of them used in other functional languages are just names of values. As such, their values never change. For example, in ML, names are bound to values with the `val` declaration, whose form is exemplified in the following:

```ml
val cost = quantity * price;
```

If `cost` appears on the left side of a subsequent `val` declaration, that declaration creates a new version of the name `cost`, which has no relationship with the previous version, which is then hidden.

F# has a somewhat similar declaration that uses the `let` reserved word. The difference between F#'s `let` and ML's `val` is that `let` creates a new scope,
whereas `val` does not. In fact, `val` declarations are often nested in `let` constructs in ML. `let` and `val` are further discussed in Chapter 15.

### 7.8 Mixed-Mode Assignment

Mixed-mode expressions were discussed in Section 7.4.1. Frequently, assignment statements also are mixed mode. The design question is: Does the type of the expression have to be the same as the type of the variable being assigned, or can coercion be used in some cases of type mismatch?

C, C++, and Perl use coercion rules for mixed-mode assignment that are similar to those they use for mixed-mode expressions; that is, many of the possible type mixes are legal, with coercion freely applied.\(^7\)

In a clear departure from C++, Java and C# allow mixed-mode assignment only if the required coercion is widening.\(^8\) So, an `int` value can be assigned to a `float` variable, but not vice versa. Disallowing half of the possible mixed-mode assignments is a simple but effective way to increase the reliability of Java and C#, relative to C and C++.

Of course, in functional languages, where assignments are just used to name values, there is no such thing as a mixed-mode assignment.

---

**SUMMARY**

Expressions consist of constants, variables, parentheses, function calls, and operators. Assignment statements include target variables, assignment operators, and expressions.

The semantics of an expression is determined in large part by the order of evaluation of operators. The associativity and precedence rules for operators in the expressions of a language determine the order of operator evaluation in those expressions. Operand evaluation order is important if functional side effects are possible. Type conversions can be widening or narrowing. Some narrowing conversions produce erroneous values. Implicit type conversions, or coercions, in expressions are common, although they eliminate the error-detection benefit of type checking, thus lowering reliability.

Assignment statements have appeared in a wide variety of forms, including conditional targets, assigning operators, and list assignments.

---

\(^7\) Note that in Python and Ruby, types are associated with objects, not variables, so there is no such thing as mixed-mode assignment in those languages.

\(^8\) Not quite true: If an integer literal, which the compiler by default assigns the type `int`, is assigned to a `char`, `byte`, or `short` variable and the literal is in the range of the type of the variable, the `int` value is coerced to the type of the variable in a narrowing conversion. This narrowing conversion cannot result in an error.
**Review Questions**

1. Define *operator precedence* and *operator associativity*.
2. What is a unary operator?
3. What is an infix operator?
4. What operator usually has left associativity?
5. When do we call operators “adjacent”? 
6. What associativity rules are used by Java?
7. What is the difference between the way exponentiation operators are implemented in Fortran and Ruby?
8. How do parentheses affect the precedence rule?
9. How is a Lisp statement declared?
10. Give a solution to the problem of operand evaluation order and side effects.
11. What is an overloaded operator?
12. Define *narrowing* and *widening conversions*.
13. In JavaScript, what is the difference between `==` and `===`?
14. What is a mixed-mode expression?
15. How is referential transparency related to functional side effects?
16. What are the advantages of referential transparency?
17. How does operand evaluation order interact with functional side effects?
18. What is short-circuit evaluation?
19. Name a language that always does short-circuit evaluation of Boolean expressions. Name one that never does it.
20. How does C support relational and Boolean expressions?
21. What is the purpose of a compound assignment operator?
22. What is the associativity of C’s unary arithmetic operators?
23. What is one possible disadvantage of treating the assignment operator as if it were an arithmetic operator?
24. What two languages include multiple assignments?
25. What mixed-mode assignments are allowed in Java?
26. What mixed-mode assignments are allowed in ML?
27. What is a cast?

**Problem Set**

1. When might you want the compiler to ignore type differences in an expression?
2. What will happen if the associativity rules of a language are changed?
3. Do you think the elimination of explicit type conversion in your favorite language would be beneficial? Why or why not?

4. Would it be a good idea to eliminate all operator precedence rules and require parentheses to show the desired precedence in expressions? Why or why not?

5. Should C’s macros be included in other languages (that do not already have them)? Why or why not?

6. Should C’s ternary operator be included in other languages (that do not already have them)? Why or why not?

7. Describe a situation in which the suboperator in a programming language would not be commutative.

8. Describe a situation in which the add operator in a programming language would not be associative.

9. Assume the following rules of associativity and precedence for expressions:

<table>
<thead>
<tr>
<th>Precedence</th>
<th>Highest</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>*, /, not</td>
</tr>
<tr>
<td></td>
<td>+, −, &amp;, mod</td>
</tr>
<tr>
<td></td>
<td>- (unary)</td>
</tr>
<tr>
<td></td>
<td>=, /=, &lt;, &lt;=, &gt;=, &gt;</td>
</tr>
<tr>
<td></td>
<td>and</td>
</tr>
</tbody>
</table>

   | Lowest           | or, xor          |

   | Associativity    | Left to right    |

Show the order of evaluation of the following expressions by parenthesizing all subexpressions and placing a superscript on the right parenthesis to indicate order. For example, for the expression

\[ a + b \times c + d \]

the order of evaluation would be represented as

\[ ((a + (b \times c)^1)^2 + d)^3 \]

a. \[ a \times b - 1 + c \]

b. \[ a \times (b - 1) / c \mod d \]

c. \[ (a - b) / c & (d \times e / a - 3) \]

d. \[ -a \text{ or } c = d \text{ and } e \]

e. \[ a > b \text{ xor } c \text{ or } d \leq 17 \]

f. \[ -a + b \]

10. Show the order of evaluation of the expressions of Problem 9, assuming that there are no precedence rules and all operators associate right to left.
11. Write a BNF description of the precedence and associativity rules defined for the expressions in Problem 9. Assume the only operands are the names a, b, c, d, and e.

12. Using the grammar of Problem 11, draw parse trees for the expressions of Problem 9.

13. Let the function fun be defined as

```c
int fun(int*k) {
    *k += 4;
    return 3 * (*k) - 1;
}
```

Suppose fun is used in a program as follows:

```c
void main() {
    int i = 10, j = 10, sum1, sum2;
    sum1 = (i / 2) + fun(&i);
    sum2 = fun(&j) + (j / 2);
}
```

What are the values of sum1 and sum2
a. operands in the expressions are evaluated left to right?
b. operands in the expressions are evaluated right to left?

14. What is your primary argument against (or for) the operator precedence rules of APL?

15. Explain why operator overloading increases the semantic capability of a language.

16. For some language of your choice, make up a list of operator symbols that could be used to eliminate all operator overloading.

17. Should automatic type conversion be included in a programming language? Why or why not?

18. Should an optimizing compiler for C or C++ be allowed to change the order of subexpressions in a Boolean expression? Why or why not?

19. Consider the following C program:

```c
int fun(int *i) {
    *i += 5;
    return 4;
}
```
Chapter 7  Expressions and Assignment Statements

void main() {
    int x = 3;
    x = x + fun(&x);
}

What is the value of $x$ after the assignment statement in main, assuming

a. operands are evaluated left to right.
b. operands are evaluated right to left.

20. Why does Java specify that operands in expressions are all evaluated in left-to-right order?

21. Explain how the coercion rules of a language affect its error detection.

PROGRAMMING EXERCISES

1. Run the code given in Problem 13 (in the Problem Set) on some system that supports C to determine the values of $sum1$ and $sum2$. Explain the results.

2. Rewrite the program of Programming Exercise 1 in C++, Java, and C#, run them, and compare the results.

3. Write a test program in your favorite language that determines and outputs the precedence and associativity of its arithmetic and Boolean operators.

4. Write a Java program that exposes Java’s rule for operand evaluation order when one of the operands is a method call.

5. Repeat Programming Exercise 4 with C++.

6. Repeat Programming Exercise 4 with C#.

7. Write a program in either C++, Java, or C# that illustrates the order of evaluation of expressions used as actual parameters to a method.

8. Write a C program that has the following statements:

```c
int a, b;
a = 10;
b = a + fun();
printf("With the function call on the right, ");
printf(" b is: %d\n", b);
a = 10;
b = fun() + a;
printf("With the function call on the left, ");
printf(" b is: %d\n", b);
```

and define fun to add 10 to a. Explain the results.

9. Write a program in either Java, C++, or C# that performs a large number of floating-point operations and an equal number of integer operations and compare the time required.
Statement-Level Control Structures

8.1 Introduction
8.2 Selection Statements
8.3 Iterative Statements
8.4 Unconditional Branching
8.5 Guarded Commands
8.6 Conclusions
The flow of control, or execution sequence, in a program can be examined at several levels. In Chapter 7, the flow of control within expressions, which is governed by operator associativity and precedence rules, was discussed. At the highest level is the flow of control among program units, which is discussed in Chapters 9 and 13. Between these two extremes is the important issue of the flow of control among statements, which is the subject of this chapter.

We begin by giving an overview of the evolution of control statements. This topic is followed by a thorough examination of selection statements, both those for two-way and those for multiple selection. Next, we discuss the variety of looping statements that have been developed and used in programming languages. Next, we take a brief look at the problems associated with unconditional branch statements. Finally, we describe the guarded command control statements.

### 8.1 Introduction

Computations in imperative-language programs are accomplished by evaluating expressions and assigning the resulting values to variables. However, there are few useful programs that consist entirely of assignment statements. At least two additional linguistic mechanisms are necessary to make the computations in programs flexible and powerful: some means of selecting among alternative control flow paths (of statement execution) and some means of causing the repeated execution of statements or sequences of statements. Statements that provide these kinds of capabilities are called **control statements**.

Computations in functional programming languages are accomplished by evaluating expressions and applying functions to given parameters. Furthermore, the flow of execution among the expressions and functions is controlled by other expressions and functions, although some of them are similar to the control statements in the imperative languages.

The control statements of the first successful programming language, Fortran, were, in effect, designed by the architects of the IBM 704. All were directly related to machine language instructions, so their capabilities were more the result of instruction design rather than language design. At the time, little was known about the difficulty of programming, and, as a result, the control statements of Fortran in the mid-1950s were thought to be entirely adequate. By today’s standards, however, they are considered seriously lacking.

A great deal of research and discussion was devoted to control statements in the 10 years between the mid-1960s and the mid-1970s. One of the primary conclusions of these efforts was that, although a single control statement (a selectable goto) is minimally sufficient, a language that is designed not to include a goto needs only a small number of different control statements. In fact, it was proven that all algorithms that can be expressed by flowcharts can be coded in a programming language with only two control statements: one for choosing between two control flow paths and one for logically controlled iterations (Böhm and Jacopini, 1966). An important result of this is that the unconditional branch statement is superfluous—potentially useful but
nonessential. This fact, combined with the practical problems of using unconditional branches, or gotos, led to a great deal of debate about the goto, as will be discussed in Section 8.4.

Programmers care less about the results of theoretical research on control statements than they do about writability and readability. All languages that have been widely used include more control statements than the two that are minimally required, because writability is enhanced by a larger number and wider variety of control statements. For example, rather than requiring the use of a logically controlled loop statement for all loops, it is easier to write programs when a counter-controlled loop statement can be used to build loops that are naturally controlled by a counter. The primary factor that restricts the number of control statements in a language is readability, because the presence of a large number of statement forms demands that program readers learn a larger language. Recall that few people learn all of the statements of a relatively large language; instead, they learn the subset they choose to use, which is often a different subset from that used by the programmer who wrote the program they are trying to read. On the other hand, too few control statements can require the use of lower-level statements, such as the goto, which also makes programs less readable.

The question as to the best collection of control statements to provide the required capabilities and the desired writability has been widely debated. It is essentially a question of how much a language should be expanded to increase its writability at the expense of its simplicity, size, and readability.

A control structure is a control statement and the collection of statements whose execution it controls.

There is only one design issue that is relevant to all of the selection and iteration control statements: Should the control structure have multiple entries? All selection and iteration constructs control the execution of code segments, and the question is whether the execution of those code segments always begins with the first statement in the segment. It is now generally believed that multiple entries add little to the flexibility of a control statement, relative to the decrease in readability caused by the increased complexity. Note that multiple entries are possible only in languages that include gotos and statement labels.

At this point, the reader might wonder why multiple exits from control structures are not considered a design issue. The reason is that all programming languages allow some form of multiple exits from control structures, the rationale being as follows: If all exits from a control structure are restricted to transferring control to the first statement following the structure, where control would flow if the control structure had no explicit exit, there is no harm to readability and also no danger. However, if an exit can have an unrestricted target and therefore can result in a transfer of control to anywhere in the program unit that contains the control structure, the harm to readability is the same as for a goto statement anywhere else in a program. Languages that have a goto statement allow it to appear anywhere, including in a control structure. Therefore, the issue is the inclusion of a goto, not whether multiple exits from control expressions are allowed.
8.2 Selection Statements

A selection statement provides the means of choosing between two or more execution paths in a program. Such statements are fundamental and essential parts of all programming languages, as was proven by Böhm and Jacopini.

Selection statements fall into two general categories: two-way and n-way, or multiple selection. Two-way selection statements are discussed in Section 8.2.1; multiple-selection statements are covered in Section 8.2.2.

8.2.1 Two-Way Selection Statements

Although the two-way selection statements of contemporary imperative languages are quite similar, there are some variations in their designs. The general form of a two-way selector is as follows:

```lang
if control_expression
    then clause
else clause
```

8.2.1.1 Design Issues

The design issues for two-way selectors can be summarized as follows:

- What is the form and type of the expression that controls the selection?
- How are the then and else clauses specified?
- How should the meaning of nested selectors be specified?

8.2.1.2 The Control Expression

Control expressions are specified in parentheses if the `then` reserved word (or some other syntactic marker) is not used to introduce the then clause. In those cases where the `then` reserved word (or alternative marker) is used, there is less need for the parentheses, so they are often omitted, as in Ruby.

In C89, which did not have a Boolean data type, arithmetic expressions were used as control expressions. This can also be done in Python, C99, and C++. However, in those languages either arithmetic or Boolean expressions can be used. In other contemporary languages, only Boolean expressions can be used for control expressions.

8.2.1.3 Clause Form

In many languages, the then and else clauses appear as either single statements or compound statements. One variation of this is Perl, in which all then and else clauses must be compound statements, even if they have only one statement. Many languages use braces to form compound statements, which serve
as the bodies of then and else clauses. In Python and Ruby, the then and else clauses are statement sequences, rather than compound statements. The complete selection statement is terminated in these languages with a reserved word. Python uses indentation to specify compound statements. For example,

```python
if x > y :
    x = y
    print "case 1"
```

All statements equally indented are included in the compound statement. Notice that rather than `then`, a colon is used to introduce the then clause in Python.

The variations in clause form have implications for the specification of the meaning of nested selectors, as discussed in the next subsection.

### 8.2.1.4 Nesting Selectors

Recall that in Chapter 3, we discussed the problem of syntactic ambiguity of a straightforward grammar for a two-way selector statement. That ambiguous grammar was as follows:

```
<if_stmt> → if <logic_expr> then <stmt>
            | if <logic_expr> then <stmt> else <stmt>
```

The issue was that when a selection statement is nested in the then clause of a selection statement, it is not clear to which if an else clause should be associated. This problem is reflected in the semantics of selection statements. Consider the following Java-like code:

```java
if (sum == 0)
    if (count == 0)
        result = 0;
else
    result = 1;
```

This statement can be interpreted in two different ways, depending on whether the else clause is matched with the first then clause or the second. Notice that the indentation seems to indicate that the else clause belongs with the first then clause. However, with the exceptions of Python and F#, indentation has no effect on semantics in contemporary languages and is therefore ignored by their compilers.

The crux of the problem in this example is that the else clause follows two then clauses with no intervening else clause, and there is no syntactic indicator to specify a matching of the else clause to one of the then clauses. In Java, as in

---
1. The statement following the compound statement must have the same indentation as the `if`.
many other imperative languages, the static semantics of the language specify that the else clause is always paired with the nearest previous unpaired then clause. A static semantics rule, rather than a syntactic entity, is used to provide the disambiguation. So, in the example, the else clause would be paired with the second then clause. The disadvantage of using a rule rather than some syntactic entity is that although the programmer may have meant the else clause to be the alternative to the first then clause and the compiler found the structure syntactically correct, its semantics is the opposite. To force the alternative semantics in Java, the inner if is put in a compound, as in

```java
if (sum == 0) {
    if (count == 0)
        result = 0;
} else
    result = 1;
```

C, C++, and C# have the same problem as Java with selection statement nesting. Because Perl requires that all then and else clauses be compound, it does not. In Perl, the previous code would be written as follows:

```perl
if (sum == 0) {
    if (count == 0) {
        result = 0;
    }
} else {
    result = 1;
}
```

If the alternative semantics were needed, it would be

```java
if (sum == 0) {
    if (count == 0) {
        result = 0;
    }
} else {
    result = 1;
}
```

Another way to avoid the issue of nested selection statements is to use an alternative means of forming compound statements. Consider the syntactic structure of the Java if statement. The then clause follows the control expression and the else clause is introduced by the reserved word else. When the then clause is a single statement and the else clause is present, although there is no need to mark the end, the else reserved word in fact marks the end of the then clause. When the then clause is a compound, it is terminated by a right brace. However, if the last clause in an if, whether then or else, is not a compound, there is no syntactic
entity to mark the end of the whole selection statement. The use of a special word for this purpose resolves the question of the semantics of nested selectors and also adds to the readability of the statement. This is the design of the selection statement in Ruby and Lua. For example, consider the following Ruby statement:

```ruby
if a > b then
  sum = sum + a
  account = account + 1
else
  sum = sum + b
  bcount = bcount + 1
end
```

The design of this statement is more regular than that of the selection statements of the C-based languages, because the form is the same regardless of the number of statements in the then and else clauses. (This is also true for Perl.) Recall that in Ruby, the then and else clauses consist of statement sequences rather than compound statements. The first interpretation of the selector example at the beginning of this section, in which the else clause is matched to the nested `if`, can be written in Ruby as follows:

```ruby
if sum == 0 then
  if count == 0 then
    result = 0
  else
    result = 1
  end
end
```

Because the `end` reserved word closes the nested `if`, it is clear that the else clause is matched to the inner then clause.

The second interpretation of the selection statement at the beginning of this section, in which the else clause is matched to the outer `if`, can be written in Ruby as follows:

```ruby
if sum == 0 then
  if count == 0 then
    result = 0
  end
else
  result = 1
end
```

The following statement, written in Python, is semantically equivalent to the last Ruby statement above:

```python
if sum == 0:
  if count == 0:
    result = 0
else:
  result = 1
```
If the line `else:` were indented to begin in the same column as the nested `if`, the else clause would be matched with the inner `if`. ML does not have a problem with nested selectors because it does not allow else-less `if` statements.

### 8.2.1.5 Selector Expressions

In the functional languages ML, F#, and LISP, the selector is not a statement; it is an expression that results in a value. Therefore, it can appear anywhere any other expression can appear. Consider the following example selector written in F#:

```fsharp
let y =
    if x > 0 then  x
    else 2 * x;;
```

This creates the name `y` and sets it to either `x` or `2 * x`, depending on whether `x` is greater than zero.

In F#, the type of the value returned by the then clause of an `if` construct must be the same as that of the value returned by its else clause. If there is no else clause, the then clause cannot return a value of a normal type. In this case, it can only return a `unit` type, which is a special type that means no value. A `unit` type is represented in code as `()`.

### 8.2.2 Multiple-Selection Statements

The multiple-selection statement allows the selection of one of any number of statements or statement groups. It is, therefore, a generalization of a selector. In fact, two-way selectors can be built with a multiple selector.

The need to choose from among more than two control paths in programs is common. Although a multiple selector can be built from two-way selectors and gotos, the resulting structures are cumbersome, unreliable, and difficult to write and read. Therefore, the need for a special structure is clear.

### 8.2.2.1 Design Issues

Some of the design issues for multiple selectors are similar to some of those for two-way selectors. For example, one issue is the question of the type of expression on which the selector is based. In this case, the range of possibilities is larger, in part because the number of possible selections is larger. A two-way selector needs an expression with only two possible values. Another issue is whether single statements, compound statements, or statement sequences may be selected. Next, there is the question of whether only a single selectable segment can be executed when the statement is executed. This is not an issue for two-way selectors, because they always allow only one of the clauses to be on a control path during one execution. As we shall see, the resolution of this issue for multiple selectors is a trade-off between reliability and flexibility. Another
issue is the form of the case value specifications. Finally, there is the issue of what should result from the selector expression evaluating to a value that does not select one of the segments. (Such a value would be unrepresented among the selectable segments.) The choice here is between simply disallowing the situation from arising and having the statement do nothing at all when it does arise.

The following is a summary of these design issues:

- What is the form and type of the expression that controls the selection?
- How are the selectable segments specified?
- Is execution flow through the structure restricted to include just a single selectable segment?
- How are the case values specified?
- How should unrepresented selector expression values be handled, if at all?

### 8.2.2.2 Examples of Multiple Selectors

The C multiple-selector statement, `switch`, which is also part of C++, Java, and JavaScript, is a relatively primitive design. Its general form is

```c
switch (expression) {
    case constant_expression_1: statement_1;
    . . .
    case constant_n: statement_n;
    [default: statement_{n+1}]
}
```

where the control expression and the constant expressions are some discrete type. This includes integer types, as well as characters and enumeration types. The selectable statements can be statement sequences, compound statements, or blocks. The optional `default` segment is for unrepresented values of the control expression. If the value of the control expression is not represented and no default segment is present, then the statement does nothing.

The `switch` statement does not provide implicit branches at the end of its code segments. This allows control to flow through more than one selectable code segment on a single execution. Consider the following example:

```c
switch (index) {
    case 1:
    case 3: odd += 1;
        sumodd += index;
    case 2:
    case 4: even += 1;
        sumeven += index;
    default: printf("Error in switch, index = %d\n", index);
}
```
This code prints the error message on every execution. Likewise, the code for the 2 and 4 constants is executed every time the code at the 1 or 3 constants is executed. To separate these segments logically, an explicit branch must be included. The `break` statement, which is actually a restricted goto, is normally used for exiting `switch` statements. `break` transfers control to the first statement after the compound statement in which it appears.

The following `switch` statement uses `break` to restrict each execution to a single selectable segment:

```c
switch (index) {
    case 1:
    case 3:  odd += 1;
             sumodd += index;
             break;
    case 2:
    case 4:  even += 1;
             sumeven += index;
             break;
    default: printf("Error in switch, index = %d\n", index);
}
```

Occasionally, it is convenient to allow control to flow from one selectable code segment to another. For example, in the example above, the segments for the case values 1 and 2 are empty, allowing control to flow to the segments for 3 and 4, respectively. This is the reason why there are no implicit branches in the `switch` statement. The reliability problem with this design arises when the mistaken absence of a `break` statement in a segment allows control to flow to the next segment incorrectly. The designers of C’s `switch` traded a decrease in reliability for an increase in flexibility. Studies have shown, however, that the ability to have control flow from one selectable segment to another is rarely used. C’s `switch` is modeled on the multiple-selection statement in ALGOL 68, which also does not have implicit branches from selectable segments.

The C switch statement has virtually no restrictions on the placement of the case expressions, which are treated as if they were normal statement labels. This laxness can result in highly complex structure within the switch body. The following example is taken from Harbison and Steele (2002).

```c
switch (x)
    default:
        if (prime(x))
            case 2: case 3: case 5: case 7:
                process_prime(x);
        else
            case 4: case 6: case 8: case 9: case 10:
                process_composite(x);
```
This code may appear to have diabolically complex form, but it was designed for a real problem and works correctly and efficiently to solve that problem.\footnote{The problem is to call \texttt{process\_prime} when \texttt{x} is prime and \texttt{process\_composite} when \texttt{x} is not prime. The design of the switch body resulted from an attempt to optimize based on the knowledge that \texttt{x} was most often in the range of 1 to 10.}

The Java switch prevents this sort of complexity by disallowing case expressions from appearing anywhere except the top level of the body of the switch.

The C\# switch statement differs from that of its C-based predecessors in two ways. First, C\# has a static semantics rule that disallows the implicit execution of more than one segment. The rule is that every selectable segment must end with an explicit unconditional branch statement: either a \texttt{break}, which transfers control out of the \texttt{switch} statement, or a \texttt{goto}, which can transfer control to one of the selectable segments (or virtually anywhere else). For example,

```csharp
switch (value) {
    case -1:
        Negatives++;
        break;
    case 0:
        Zeros++;
        goto case 1;
    case 1:
        Positives++;
    default:
        Console.WriteLine("Error in switch \n");
}
```

Note that \texttt{Console.WriteLine} is the method for displaying strings in C#.

The other way C\#'s \texttt{switch} differs from that of its predecessors is that the control expression and the case statements can be strings in C#.

PHP's \texttt{switch} uses the syntax of C's \texttt{switch} but allows more type flexibility. The case values can be any of the PHP scalar types—string, integer, or double precision. As with C, if there is no \texttt{break} at the end of the selected segment, execution continues into the next segment.

Ruby has two forms of multiple-selection constructs, both of which are called \texttt{case expressions} and both of which yield the value of the last expression evaluated. The only version of Ruby's case expressions that is described here is semantically similar to a list of nested if statements:

```ruby
case
when Boolean_expression then expression
  . . .
when Boolean_expression then expression
[else expression]
end
```

The Java switch prevents this sort of complexity by disallowing case expressions from appearing anywhere except the top level of the body of the switch.
The semantics of this case expression is that the Boolean expressions are evaluated one at a time, top to bottom. The value of the case expression is the value of the first then expression whose Boolean expression is true. The else represents true in this statement, and the else clause is optional. For example,³

\[
\text{leap} = \text{case} \begin{array}{l}
\text{when} \ year \ % \ 400 == 0 \ \text{then} \ true \\
\text{when} \ year \ % \ 100 == 0 \ \text{then} \ false \\
\text{else} \ year \ % \ 4 == 0 \\
\end{array} \text{end}
\]

This case expression evaluates to true if \textit{year} is a leap year.

The other Ruby case expression form is similar to the switch of Java. Perl, Python, and Lua do not have multiple-selection statements.

### 8.2.2.3 Implementing Multiple Selection Structures

A multiple selection statement is essentially an n-way branch to segments of code, where n is the number of selectable segments. Implementing such a statement must be done with multiple conditional branch instructions. Consider again the general form of the C switch statement, with breaks:

\[
\text{switch} (\text{expression}) \{ \\
\quad \text{case constant_expression}_1; \text{statement}_1; \\
\quad \quad \text{break;} \\
\quad \quad \ldots \\
\quad \quad \text{case constant}_n; \text{statement}_n; \\
\quad \quad \quad \text{break;} \\
\quad \quad \quad [\text{default}; \text{statement}_{n+1}] \\
\}
\]

One simple translation of this statement follows:

Code to evaluate expression into t
\[
\text{goto} \ \text{branches} \\
\text{label}_1: \text{code for statement}_1 \\
\quad \text{goto} \ \text{out} \\
\quad \ldots \\
\text{label}_n: \text{code for statement}_n \\
\quad \text{goto} \ \text{out} \\
\text{default}: \text{code for statement}_{n+1} \\
\quad \text{goto} \ \text{out} \\
\text{branches: if } t = \text{constant_expression}_1 \ \text{goto} \ \text{label}_1 \\
\quad \ldots \\
\quad \quad \text{if } t = \text{constant_expression}_n \ \text{goto} \ \text{label}_n \\
\quad \text{goto} \ \text{default} \\
\text{out:}
\]

³. This example is from Thomas et al. (2013).
The code for the selectable segments precedes the branches so that the targets of the branches are all known when the branches are generated. An alternative to these coded conditional branches is to put the case values and labels in a table and use a linear search with a loop to find the correct label. This requires less space than the coded conditionals.

The use of conditional branches or a linear search on a table of cases and labels is a simple but inefficient approach that is acceptable when the number of cases is small, say less than 10. It takes an average of about half as many tests as there are cases to find the right one. For the default case to be chosen, all other cases must be tested. In statements with 10 or more cases, the low efficiency of this form is not justified by its simplicity.

When the number of cases is 10 or greater, the compiler can build a hash table of the segment labels, which would result in approximately equal (and short) times to choose any of the selectable segments. If the language allows ranges of values for case expressions, as in Ruby, the hash method is not suitable. For these situations, a binary search table of case values and segment addresses is better.

If the range of the case values is relatively small and more than half of the whole range of values is represented, an array whose indices are the case values and whose values are the segment labels can be built. Array elements whose indices are not among the represented case values are filled with the default segment's label. Then finding the correct segment label is found by array indexing, which is very fast.

Of course, choosing among these approaches is an additional burden on the compiler. In many compilers, only two different methods are available. As in other situations, determining and using the most efficient method costs more compiler time.

8.2.2.4 Multiple Selection Using if

In many situations, a switch or case statement is inadequate for multiple selection (Ruby’s case is an exception). For example, when selections must be made on the basis of a Boolean expression rather than some ordinal type, nested two-way selectors can be used to simulate a multiple selector. To alleviate the poor readability of deeply nested two-way selectors, some languages, such as Perl and Python, have been extended specifically for this use. The extension allows some of the special words to be left out. In particular, else-if sequences are replaced with a single special word, and the closing special word on the nested if is dropped. The nested selector is then called an else-if clause. Consider the following Python selector statement (note that else-if is spelled elif in Python):

```python
if count < 10 :
    bag1 = True
elif count < 100 :
    bag2 = True
elif count < 1000 :
    bag3 = True
```
which is equivalent to the following:

```python
if count < 10 :
    bag1 = True
else :
    if count < 100 :
        bag2 = True
    else :
        if count < 1000 :
            bag3 = True
        else :
            bag4 = True
```

The else-if version (the first) is the more readable of the two. Notice that this example is not easily simulated with a `switch` statement, because each selectable statement is chosen on the basis of a Boolean expression. Therefore, the else-if statement is not a redundant form of `switch`. In fact, none of the multiple selectors in contemporary languages are as general as the if-then-else-if statement. An operational semantics description of a general selector statement with else-if clauses, in which the E’s are logic expressions and the S’s are statements, is given here:

```plaintext
if E_1 goto 1
if E_2 goto 2
...
1: S_1
goto out
2: S_2
goto out
...
out: ...
```

From this description, we can see the difference between multiple selection structures and else-if statements: In a multiple selection statement, all the E’s would be restricted to comparisons between the value of a single expression and some other values.

Languages that do not include the else-if statement can use the same control structure, with only slightly more typing.

The Python example if-then-else-if statement above can be written as the Ruby `case` statement:

```ruby
case
  when count < 10 then bag1 = true
  when count < 100 then bag2 = true
  when count < 1000 then bag3 = true
end
```

Else-if statements are based on the common mathematics statement, the conditional expression.
The Scheme multiple selector, which is based on mathematical conditional expressions, is a special form function named **COND**. **COND** is a slightly generalized version of the mathematical conditional expression; it allows more than one predicate to be true at the same time. Because different mathematical conditional expressions have different numbers of parameters, **COND** does not require a fixed number of actual parameters. Each parameter to **COND** is a pair of expressions in which the first is a predicate (it evaluates to either #T or #F).

The general form of **COND** is

```
(COND
  (predicate1 expression1)
  (predicate2 expression2)
  ...
  (predicaten expressionn)
  [ (ELSE expressionn+1) ]
)
```

where the **ELSE** clause is optional.

The semantics of **COND** is as follows: The predicates of the parameters are evaluated one at a time, in order from the first, until one evaluates to #T. The expression that follows the first predicate that is found to be #T is then evaluated and its value is returned as the value of **COND**. If none of the predicates is true and there is an **ELSE**, its expression is evaluated and the value is returned. If none of the predicates is true and there is no **ELSE**, the value of **COND** is unspecified. Therefore, all **CONDs** should include an **ELSE**.

Consider the following example call to **COND**:

```
(COND
  ((> x y) "x is greater than y")
  ((< x y) "y is greater than x")
  (ELSE "x and y are equal")
)
```

Note that string literals evaluate to themselves, so that when this call to **COND** is evaluated, it produces a string result.

### 8.3 Iterative Statements

An **iterative statement** is one that causes a statement or collection of statements to be executed zero, one, or more times. An iterative statement is often called a **loop**. Every programming language from Plankalkül on has included some method of repeating the execution of segments of code. Iteration is the very essence of the power of the computer. If some means of repetitive execution of a statement or collection of statements were not possible, programmers would be required to state every action in sequence; useful programs would be
huge and inflexible and take unacceptably large amounts of time to write and mammoth amounts of memory to store.

The first iterative statements in programming languages were directly related to arrays. This resulted from the fact that in the earliest years of computers, computing was largely numerical in nature, frequently using loops to process data in arrays.

Several categories of iteration control statements have been developed. The primary categories are defined by how designers answered two basic design questions:

- How is the iteration controlled?
- Where should the control mechanism appear in the loop statement?

The primary possibilities for iteration control are logical, counting, or a combination of the two. The main choices for the location of the control mechanism are the top of the loop or the bottom of the loop. Top and bottom here are logical, rather than physical, denotations. The issue is not the physical placement of the control mechanism; rather, it is whether the mechanism is executed and affects control before or after execution of the statement’s body. A third option, which allows the user to decide where to put the control, at the top, at the bottom, or even within the controlled segment, is discussed in Section 8.3.3.

The body of an iterative statement is the collection of statements whose execution is controlled by the iteration statement. We use the term pretest to mean that the test for loop completion occurs before the loop body is executed and posttest to mean that it occurs after the loop body is executed. The iteration statement and the associated loop body together form an iteration statement.

### 8.3.1 Counter-Controlled Loops

A counting iterative control statement has a variable, called the loop variable, in which the count value is maintained. It also includes some means of specifying the initial and terminal values of the loop variable, and the difference between sequential loop variable values, often called the stepsize. The initial, terminal, and stepsize specifications of a loop are called the loop parameters.

Although logically controlled loops are more general than counter-controlled loops, they are not necessarily more commonly used. Because counter-controlled loops are more complex, their design is more demanding.

Counter-controlled loops are sometimes supported by machine instructions designed for that purpose. Unfortunately, machine architecture can outlive the prevailing approaches to programming at the time of the architecture design. For example, VAX computers have a very convenient instruction for the implementation of posttest counter-controlled loops, which Fortran had at the time of the design of the VAX (mid-1970s). But Fortran no longer had such a loop by the time VAX computers became widely used (it had been replaced by a pretest
loop). Furthermore, no other widely used language of the time had a posttest counting loop.

### 8.3.1.1 Design Issues

There are many design issues for iterative counter-controlled statements. The nature of the loop variable and the loop parameters provide a number of design issues. The type of the loop variable and that of the loop parameters obviously should be the same or at least compatible, but what types should be allowed? One apparent choice is integer, but what about enumeration, character, and floating-point types? Another question is whether the loop variable is a normal variable, in terms of scope, or whether it should have some special scope. Allowing the user to change the loop variable or the loop parameters within the loop can lead to code that is very difficult to understand, so another question is whether the additional flexibility that might be gained by allowing such changes is worth that additional complexity. A similar question arises about the number of times and the specific time when the loop parameters are evaluated: If they are evaluated just once, loops are simple but less flexible.

The following is a summary of these design issues:

- What are the type and scope of the loop variable?
- Should it be legal for the loop variable or loop parameters to be changed in the loop, and if so, does the change affect loop control?
- Should the loop parameters be evaluated only once, or once for every iteration?

### 8.3.1.2 The for Statement of the C-Based Languages

The general form of C’s `for` statement is

```c
for (expression_1; expression_2; expression_3)
  loop body
```

The loop body can be a single statement, a compound statement, or a null statement.

Because assignment statements in C produce results and thus can be considered expressions, the expressions in a `for` statement are often assignment statements. The first expression is for initialization and is evaluated only once, when the `for` statement execution begins. The second expression is the loop control and is evaluated before each execution of the loop body. As is usual in C, a zero value means false and all nonzero values mean true. Therefore, if the value of the second expression is zero, the `for` is terminated; otherwise, the loop body statements are executed. In C99, the expression also could be a Boolean type. A C99 Boolean type stores only the values 0 or 1. The last expression in the `for` is executed after each execution of the loop body. It is often used
to increment the loop counter. An operational semantics description of the C
for statement is shown next. Because C expressions can be used as statements,
expression evaluations are shown as statements.

    expression_1
loop:    
     if expression_2 = 0 goto out
 [loop body]
    expression_3
    goto loop
out: . . .

Following is an example of a skeletal C for statement:

    for (count = 1; count <= 10; count++)
    . . .
}

All of the expressions of C’s for are optional. An absent second expression is
considered true, so a for without one is potentially an infinite loop. If the first
and/or third expressions are absent, no assumptions are made. For example, if
the first expression is absent, it simply means that no initialization takes place.

    Note that C’s for need not count. It can easily model counting and logical
loop structures, as demonstrated in the next section.

    The C for design choices are the following: There are no explicit loop
variables or loop parameters. All involved variables can be changed in the loop
body. The expressions are evaluated in the order stated previously. Although it
can create havoc, it is legal to branch into a C for loop body.

    C’s for is one of the most flexible, because each of the expressions can
comprise multiple expressions, which in turn allow multiple loop variables that
can be of any type. When multiple expressions are used in a single expression
of a for statement, they are separated by commas. All C statements have val-
ues, and this form of multiple expression is no exception. The value of such a
multiple expression is the value of the last component.

    Consider the following for statement:

    for (count1 = 0, count2 = 1.0;
    count1 <= 10 && count2 <= 100.0;
    sum = ++count1 + count2, count2 *= 2.5);

The operational semantics description of this is

    count1 = 0
    count2 = 1.0
loop:
    if count1 > 10 goto out
    if count2 > 100.0 goto out
    count1 = count1 + 1
8.3 Iterative Statements

The example C for statement does not need and thus does not have a loop body. All the desired actions are part of the for statement itself, rather than in its body. The first and third expressions are multiple statements. In both of these cases, the whole expression is evaluated, but the resulting value is not used in the loop control.

The for statement of C99 and C++ differs from that of earlier versions of C in two ways. First, in addition to an arithmetic expression, it can use a Boolean expression for loop control. Second, the first expression can include variable definitions. For example,

```c
for (int count = 0; count < len; count++) { /* ... */ }
```

The scope of a variable defined in the for statement is from its definition to the end of the loop body.

The for statement of Java and C# is like that of C++, except that the loop control expression is restricted to boolean.

In all of the C-based languages, the last two loop parameters are evaluated with every iteration. Furthermore, variables that appear in the loop parameter expression can be changed in the loop body. Therefore, these loops can be complex and potentially unreliable.

### 8.3.1.3 The for Statement of Python

The general form of Python’s for is

```python
for loop_variable in object:
    - loop body
[else:
    - else clause]
```

The loop variable is assigned the value in the object, which is often a range, one for each execution of the loop body. The else clause, when present, is executed if the loop terminates normally.

Consider the following example:

```python
for count in [2, 4, 6]:
    print count
```

produces

```
2
4
6
```
For most simple counting loops in Python, the `range` function is used. `range` takes one, two, or three parameters. The following examples demonstrate the actions of `range`:

- `range(5)` returns `[0, 1, 2, 3, 4]`
- `range(2, 7)` returns `[2, 3, 4, 5, 6]`
- `range(0, 8, 2)` returns `[0, 2, 4, 6]`

Note that `range` never returns the highest value in a given parameter range.

### 8.3.1.4 Counter-Controlled Loops in Functional Languages

Counter-controlled loops in imperative languages use a counter variable, but such variables do not exist in pure functional languages. Rather than iteration to control repetition, functional languages use recursion. Rather than a statement, functional languages use a recursive function. Counting loops can be simulated in functional languages as follows: The counter can be a parameter for a function that repeatedly executes the loop body, which can be specified in a second function sent to the loop function as a parameter. So, such a loop function takes the body function and the number of repetitions as parameters.

The general form of an F# function for simulating counting loops, named `forLoop` in this case, is as follows:

```fsharp
let rec forLoop loopBody reps =
    if reps <= 0 then
        ()
    else
        loopBody()
        forLoop loopBody, (reps - 1);;
```

In this function, the parameter `loopBody` is the function with the body of the loop and the parameter `reps` is the number of repetitions. The reserved word `rec` appears before the name of the function to indicate that it is recursive. The empty parentheses do nothing; they are there because in F# an empty statement is illegal and every `if` must have an else clause.

### 8.3.2 Logically Controlled Loops

In many cases, collections of statements must be repeatedly executed, but the repetition control is based on a Boolean expression rather than a counter. For these situations, a logically controlled loop is convenient. Actually, logically controlled loops are more general than counter-controlled loops. Every counting loop can be built with a logical loop, but the reverse is not true. Also, recall that only selection and logical loops are essential to express the control structure of any flowchart.

#### 8.3.2.1 Design Issues

Because they are much simpler than counter-controlled loops, logically controlled loops have fewer design issues.
• Should the control be pretest or posttest?
• Should the logically controlled loop be a special form of a counting loop or a separate statement?

8.3.2.2 Examples

The C-based programming languages include both pretest and posttest logically controlled loops that are not special forms of their counter-controlled iterative statements. The pretest and posttest logical loops have the following forms:

```
while (control_expression)
  loop body

and

do
  loop body
while (control_expression);
```

These two statement forms are exemplified by the following C# code segments:

```
sum = 0;
indat = Int32.Parse(Console.ReadLine());
while (indat >= 0) {
  sum += indat;
  indat = Int32.Parse(Console.ReadLine());
}

value = Int32.Parse(Console.ReadLine());
do {
  value /= 10;
  digits ++;
} while (value > 0);
```

Note that all variables in these examples are integer type. The `ReadLine` method of the `Console` object gets a line of text from the keyboard. `Int32.Parse` finds the number in its string parameter, converts it to `int` type, and returns it.

In the pretest version of a logical loop (`while`), the statement or statement segment is executed as long as the expression evaluates to true. In the posttest version (`do`), the loop body is executed until the expression evaluates to false. In both cases, the statement can be compound. The operational semantics descriptions of those two statements follow:

```
while
  loop:
    if control_expression is false goto out
  [loop body]
```
goto loop
out: . . .

do-while
loop:
[loop body]
if control_expression is true goto loop

It is legal in both C and C++ to branch into both while and do loop bodies. The C89 version uses an arithmetic expression for control; in C99 and C++, it may be either arithmetic or Boolean.

Java's while and do statements are similar to those of C and C++, except the control expression must be boolean type, and because Java does not have a goto, the loop bodies cannot be entered anywhere except at their beginnings.

Posttest loops are infrequently useful and also can be somewhat dangerous, in the sense that programmers sometimes forget that the loop body will always be executed at least once. The syntactic design of placing a posttest control physically after the loop body, where it has its semantic effect, helps avoid such problems by making the logic clear.

A pretest logical loop can be simulated in a purely functional form with a recursive function that is similar to the one used to simulate a counting loop statement in Section 8.3.1.5. In both cases, the loop body is written as a function. Following is the general form of a simulated logical pretest loop, written in F#:

let rec whileLoop test body =
  if test() then
    body()
    whileLoop test body
  else
    ();

8.3.3 User-Located Loop Control Mechanisms

In some situations, it is convenient for a programmer to choose a location for loop control other than the top or bottom of the loop body. As a result, some languages provide this capability. A syntactic mechanism for user-located loop control can be relatively simple, so its design is not difficult. Such loops have the structure of infinite loops but include one or more user-located loop exits. Perhaps the most interesting question is whether a single loop or several nested loops can be exited. The design issues for such a mechanism are the following:

- Should the conditional mechanism be an integral part of the exit?
- Should only one loop body be exited, or can enclosing loops also be exited?

C, C++, Python, Ruby, and C# have unconditional unlabeled exits (break). Java and Perl have unconditional labeled exits (break in Java, last in Perl).
Following is an example of nested loops in Java, in which there is a break out of the outer loop from the nested loop:

```java
outerLoop:
  for (row = 0; row < numRows; row++)
    for (col = 0; col < numCols; col++) {
      sum += mat[row][col];
      if (sum > 1000.0)
        break outerLoop;
    }
```

C, C++, and Python include an unlabeled control statement, `continue`, that transfers control to the control mechanism of the smallest enclosing loop. This is not an exit but rather a way to skip the rest of the loop statements on the current iteration without terminating the loop construct. For example, consider the following:

```java
while (sum < 1000) {
  getnext(value);
  if (value < 0) continue;
  sum += value;
}
```

A negative value causes the assignment statement to be skipped, and control is transferred instead to the conditional at the top of the loop. On the other hand, in

```java
while (sum < 1000) {
  getnext(value);
  if (value < 0) break;
  sum += value;
}
```

a negative value terminates the loop.

Both `last` and `break` provide for multiple exits from loops, which may seem to be somewhat of a hindrance to readability. However, unusual conditions that require loop termination are so common that such a statement is justified. Furthermore, readability is not seriously harmed, because the target of all such loop exits is the first statement after the loop (or an enclosing loop) rather than just anywhere in the program. Finally, the alternative of using multiple breaks to leave more than one level of loops is even more detrimental to readability.

The motivation for user-located loop exits is simple: They fulfill a common need for goto statements using a highly restricted branch statement. The target of a goto can be many places in the program, both above and below the goto itself. However, the targets of user-located loop exits must be below the exit and can only follow immediately at the end of a compound statement.
8.3.4 Iteration Based on Data Structures

A general data-based iteration statement uses a user-defined data structure and a user-defined function (the iterator) to go through the structure’s elements. The iterator is called at the beginning of each iteration, and each time it is called, the iterator returns an element from a particular data structure in some specific order. For example, suppose a program has a user-defined binary tree of data nodes, and the data in each node must be processed in some particular order. A user-defined iteration statement for the tree would successively set the loop variable to point to the nodes in the tree, one for each iteration. The initial execution of the user-defined iteration statement needs to issue a special call to the iterator to get the first tree element. The iterator must always remember which node it presented last so that it visits all nodes without visiting any node more than once. So an iterator must be history sensitive. A user-defined iteration statement terminates when the iterator fails to find more elements.

The for statement of the C-based languages, because of its great flexibility, can be used to simulate a user-defined iteration statement. Once again, suppose the nodes of a binary tree are to be processed. If the tree root is pointed to by a variable named root, and if traverse is a function that sets its parameter to point to the next element of a tree in the desired order, the following could be used:

```c
for (ptr = root; ptr != null; ptr = traverse(ptr)) {
    . . .
}
```

In this statement, traverse is the iterator.

Predefined iterators are used to provide iterative access to PHP’s unique arrays. The current pointer points at the element last accessed through iteration. The next iterator moves current to the next element in the array. The prev iterator moves current to the previous element. current can be set or reset to the array’s first element with the reset operator. The following code displays all of the elements of an array of numbers $list:

```c
reset $list;
p1rint ("First number: " + current($list) + "");
while ($current_value = next($list))
    print ("Next number: " + $current_value + "<br \>");
```

User-defined iteration statements are more important in object-oriented programming than they were in earlier software development paradigms, because users of object-oriented programming routinely use abstract data types for data structures, especially collections. In such cases, a user-defined iteration statement and its iterator must be provided by the author of the data abstraction because the representation of the objects of the type is not known to the user.

An enhanced version of the for statement was added to Java in Java 5.0. This statement simplifies iterating through the values in an array or objects
in a collection that implements the `Iterable` interface. (All of the predefined
generic collections in Java implement `Iterable`.) For example, if we had an
`ArrayList` collection named `myList` of strings, the following statement
would iterate through all of its elements, setting each to `myElement`:

```java
for (String myElement : myList) { ... }
```

This new statement is referred to as “foreach,” although its reserved word is `for`.

C# and F# (and the other .NET languages) also have generic library classes
for collections. For example, there are generic collection classes for lists, which
are dynamic length arrays, stacks, queues, and dictionaries (hash table). All of
these predefined generic collections have built-in iterators that are used implicit-
ly with the `foreach` statement. Furthermore, users can define their own col-
lections and write their own iterators, which can implement the `IEnumerator`
interface, which enables the use of `foreach` on these collections.

For example, consider the following C# code:

```csharp
List<String> names = new List<String>();
names.Add("Bob");
names.Add("Carol");
names.Add("Alice");
...
foreach (String name in names)
    Console.WriteLine(name);
```

In Ruby, a block is a sequence of code, delimited by either braces or the `do` and `end` reserved words. Blocks can be used with specially written methods
to create many useful constructs, including iterators for data structures. This
construct consists of a method call followed by a block. A block is actually
an anonymous method that is sent to the method (whose call precedes it) as
a parameter. The called method can then call the block, which can produce
output or objects.

Ruby predefines several iterator methods, such as `times` and `upto` for
counter-controlled loops, and `each` for simple iterations of arrays and hashes.
For example, consider the following example of using `times`:

```ruby
>> 4.times {puts "Hey!"}
Hey!
Hey!
Hey!
Hey!
Hey!
=> 4
```

---

4. An `ArrayList` is a predefined generic collection that is actually a dynamic-length array
of whatever type it is declared to store.
Note that >> is the prompt of the interactive Ruby interpreter and => is used to indicate the return value of the expression. The Ruby puts statement displays its parameter. In this example, the times method is sent to the object 4, with the block sent along as a parameter. The times method calls the block four times, producing the four lines of output. The destination object, 4, is the return value from times.

The most common Ruby iterator is each, which is often used to go through arrays and apply a block to each element. For this purpose, it is convenient to allow blocks to have parameters, which, if present, appear at the beginning of the block, delimited by vertical bars (|). The following example, which uses a block parameter, illustrates the use of each:

```ruby
>> list = [2, 4, 6, 8]
=> [2, 4, 6, 8]
>> list.each {|value| puts value}
2
4
6
8
=> [2, 4, 6, 8]
```

In this example, the block is called for each element of the array to which the each method is sent. The block produces the output, which is a list of the array’s elements. The return value of each is the array to which it is sent.

Instead of a counting loop, Ruby has the upto method. For example, we could have the following:

```ruby
1.upto(5) {|x| print x, " "}
```

This produces the following output:

```
1 2 3 4 5
```

Syntax that resembles a for loop in other languages could also be used, as in the following:

```ruby
for x in 1..5
    print x, " 
end
```

Ruby actually has no for statement—constructs like the above are converted by Ruby into upto method calls.

---

5. This is similar to the map functions discussed in Chapter 15.
8.4 Unconditional Branching

An **unconditional branch statement** transfers execution control to a specified location in the program. The most heated debate in language design in the late 1960s was over the issue of whether unconditional branching should be part of any high-level language, and if so, whether its use should be restricted. The unconditional branch, or goto, is the most powerful statement for controlling the flow of execution of a program’s statements. However, careless use of the goto can lead to serious problems. The goto has stunning power and great flexibility (all other control structures can be built with goto and a selector), but it is this power that makes its use dangerous. Without usage restrictions, imposed by either language design or programming standards, goto statements can make programs very difficult to read, and as a result, highly unreliable and costly to maintain.

These problems follow directly from a goto’s ability to force any program statement to follow any other in execution sequence, regardless of whether that statement precedes or follows the previously executed statement in textual order. Readability is best when the execution order of statements in a program is nearly the same as the order in which they appear—in our case, this would mean top to bottom, which is the order to which we are accustomed. Thus, restricting gotos so they can transfer control only downward in a program partially alleviates the problem. It allows gotos to transfer control around code sections in response to errors or unusual conditions but disallows their use to build any sort of loop.

A few languages have been designed without a goto—for example, Java, Python, and Ruby. However, most currently popular languages include a goto statement. Kernighan and Ritchie (1978) call the goto infinitely abusable, but it is nevertheless included in Ritchie’s language, C. The languages that have eliminated the goto have provided additional control statements, usually in the form of loop exits, to code one of the justifiable applications of the goto.

The relatively new language, C#, includes a goto, even though one of the languages on which it is based, Java, does not.
One legitimate use of C#'s goto is in the `switch` statement, as discussed in Section 8.2.2.2.

All of the loop exit statements discussed in Section 8.3.3 are actually camouflaged goto statements. They are, however, severely restricted gotos and are not harmful to readability. In fact, it can be argued that they improve readability, because to avoid their use results in convoluted and unnatural code that would be much more difficult to understand.

8.5 Guarded Commands

New and quite different forms of selection and loop structures were suggested by Dijkstra (1975). His primary motivation was to provide control statements that would support a program design methodology that ensured correctness during development rather than when verifying or testing completed programs. This methodology is described in Dijkstra (1976). Another motivation is the increased clarity in reasoning that is possible with guarded commands. Simply put, a selectable segment of a selection statement in a guarded-command statement can be considered independently of any other part of the statement, which is not true for the selection statements of the common programming languages.

Guarded commands are covered in this chapter because they are the basis for the linguistic mechanism developed later for concurrent programming in CSP (Hoare, 1978). Guarded commands are also used to define functions in Haskell, as discussed in Chapter 15.

Dijkstra's selection statement has the form

```
if <Boolean expression> -> <statement>
[ ] <Boolean expression> -> <statement>
[ ] . . .
[ ] <Boolean expression> -> <statement>
fi
```

The closing reserved word, `fi`, is the opening reserved word spelled backward. This form of closing reserved word is taken from ALGOL 68. The small blocks, called `fatbars`, are used to separate the guarded clauses and allow the clauses to be statement sequences. Each line in the selection statement, consisting of a Boolean expression (a guard) and a statement or statement sequence, is called a guarded command.

This selection statement has the appearance of a multiple selection, but its semantics is different. All of the Boolean expressions are evaluated each time the statement is reached during execution. If more than one expression is true, one of the corresponding statements can be nondeterministically chosen for execution. An implementation might always choose the statement associated with the first Boolean expression that evaluates to be true. But it may choose any statement associated with a true Boolean expression. So, the correctness
of the program cannot depend on which statement is chosen (among those associated with true Boolean expressions). If none of the Boolean expressions are true, a run-time error occurs that causes program termination. This forces the programmer to consider and list all possibilities. Consider the following example:

```plaintext
if i = 0 -> sum := sum + i
[] i > j -> sum := sum + j
[] j > i -> sum := sum + i
fi
```

If \( i = 0 \) and \( j > i \), this statement chooses nondeterministically between the first and third assignment statements. If \( i \) is equal to \( j \) and is not zero, a run-time error occurs because none of the conditions are true.

This statement can be an elegant way of allowing the programmer to state that the order of execution, in some cases, is irrelevant. For example, to find the largest of two numbers, we can use

```plaintext
if x >= y -> max := x
[] y >= x -> max := y
fi
```

This computes the desired result without overspecifying the solution. In particular, if \( x \) and \( y \) are equal, it does not matter which we assign to \( max \). This is a form of abstraction provided by the nondeterministic semantics of the statement.

Now, consider this same process coded in a traditional programming language selector:

```plaintext
if (x >= y)
    max = x;
else
    max = y;
```

This could also be coded as follows:

```plaintext
if (x > y)
    max = x;
else
    max = y;
```

There is no practical difference between these two statements. The first assigns \( x \) to \( max \) when \( x \) and \( y \) are equal; the second assigns \( y \) to \( max \) in the same circumstance. This choice between the two statements complicates the formal analysis of the code and the correctness proof of it. This is one of the reasons why guarded commands were developed by Dijkstra.
The loop structure proposed by Dijkstra has the form

\[
\text{do } \langle \text{Boolean expression} \rangle \rightarrow \langle \text{statement} \rangle \\
\text{[ ] } \langle \text{Boolean expression} \rangle \rightarrow \langle \text{statement} \rangle \\
\text{[ ] . . .} \\
\text{[ ] } \langle \text{Boolean expression} \rangle \rightarrow \langle \text{statement} \rangle \\
\text{od}
\]

The semantics of this statement is that all Boolean expressions are evaluated on each iteration. If more than one is true, one of the associated statements is nondeterministically (perhaps randomly) chosen for execution, after which the expressions are again evaluated. When all expressions are simultaneously false, the loop terminates.

Consider the following problem: Given four integer variables, \(q_1, q_2, q_3,\) and \(q_4,\) rearrange the values of the four so that \(q_1 \geq q_2 \geq q_3 \geq q_4.\) Without guarded commands, one straightforward solution is to put the four values into an array, sort the array, and then assign the values from the array back into the scalar variables \(q_1, q_2, q_3,\) and \(q_4.\) While this solution is not difficult, it requires a good deal of code, especially if the sort process must be included.

Now, consider the following code, which uses guarded commands to solve the same problem but in a more concise and elegant way.\(^6\)

\[
\text{do } q_1 > q_2 \rightarrow \text{temp} := q_1; \ q_1 := q_2; \ q_2 := \text{temp}; \\
\text{[ ] } q_2 > q_3 \rightarrow \text{temp} := q_2; \ q_2 := q_3; \ q_3 := \text{temp}; \\
\text{[ ] } q_3 > q_4 \rightarrow \text{temp} := q_3; \ q_3 := q_4; \ q_4 := \text{temp}; \\
\text{od}
\]

Dijkstra’s guarded command control statements are interesting, in part because they illustrate how the syntax and semantics of statements can have an impact on program verification and vice versa. Program verification is virtually impossible when goto statements are used. Verification is greatly simplified if (1) only logical loops and selections are used or (2) only guarded commands are used. The axiomatic semantics of guarded commands are conveniently specified (Gries, 1981). It should be obvious, however, that there is considerably increased complexity in the implementation of the guarded commands over their conventional deterministic counterparts.

### 8.6 Conclusions

We have described and discussed a variety of statement-level control structures. A brief evaluation now seems to be in order.

First, we have the theoretical result that only sequence, selection, and pretest logical loops are absolutely required to express computations (Böhm and

---

\(^6\) This code appears in a slightly different form in Dijkstra (1975).
Jacopini, 1966). This result has been used by those who wish to ban unconditional branching altogether. Of course, there are already sufficient practical problems with the goto to condemn it without also using a theoretical reason. One of the main legitimate needs for gotos—premature exits from loops—can be met with restricted branch statements, such as break.

One obvious misuse of the Böhm and Jacopini result is to argue against the inclusion of any control structures beyond selection and pretest logical loops. No widely used language has yet taken that step; furthermore, we doubt that any ever will, because of the negative effect on writability and readability. Programs written with only selection and pretest logical loops are generally less natural in structure, more complex, and therefore harder to write and more difficult to read. For example, the C# multiple selection structure is a great boost to C# writability, with no obvious negatives. Another example is the counting loop structure of many languages, especially when the statement is simple.

It is not so clear that the utility of many of the other control structures that have been proposed is worth their inclusion in languages (Ledgard and Marcotty, 1975). This question rests to a large degree on the fundamental question of whether the size of languages must be minimized. Both Wirth (1975) and Hoare (1973) strongly endorse simplicity in language design. In the case of control structures, simplicity means that only a few control statements should be in a language, and they should be simple.

The rich variety of statement-level control structures that have been invented shows the diversity of opinion among language designers. After all the invention, discussion, and evaluation, there is still no unanimity of opinion on the precise set of control statements that should be in a language. Most contemporary languages do, of course, have similar control statements, but there is still some variation in the details of their syntax and semantics. Furthermore, there is still disagreement on whether a language should include a goto; C++ and C# do, but Java and Ruby do not.

**SUMMARY**

Control statements occur in several categories: selection, multiple selection, iterative, and unconditional branching.

The switch statement of the C-based languages is representative of multiple-selection statements. The C# version eliminates the reliability problem of its predecessors by disallowing the implicit continuation from a selected segment to the following selectable segment.

A large number of different loop statements have been invented for high-level languages. C’s for statement is the most flexible iteration statement, although its flexibility leads to some reliability problems.

Most languages have exit statements for their loops; these statements take the place of one of the most common uses of goto statements.

Data-based iterators are loop statements for processing data structures, such as linked lists, hashes, and trees. The for statement of the C-based
languages allows the user to create iterators for user-defined data. The **foreach** statement of Perl and C# is a predefined iterator for standard data structures. In the contemporary object-oriented languages, iterators for collections are specified with standard interfaces, which are implemented by the designers of the collections.

Ruby includes iterators that are a special form of methods that are sent to various objects. The language predefined iteratrors for common uses, but also allows user-defined iterators.

The unconditional branch, or goto, has been part of most imperative languages. Its problems have been widely discussed and debated. The current consensus is that it should remain in most languages but that its dangers should be minimized through programming discipline.

Dijkstra’s guarded commands are alternative control statements with positive theoretical characteristics. Although they have not been adopted as the control statements of a language, part of the semantics appear in the concurrency mechanisms of CSP and the function definitions of Haskell.

### REVIEW QUESTIONS

1. Define selection statement.
2. Mention one common misuse of the Böhm and Jacopini result.
3. What is the general form of a two-way selector?
4. What mechanism does Python use to specify compound statements?
5. Why is a static semantics rule used instead of a syntactic entity in many languages?
6. Give an example of a language that does not allow else-less **if** statements.
7. Under what circumstances can arithmetic expressions be used as control expressions?
8. What is the limitation of building a multiple selector from two-way selectors and gotos?
9. What are the two general categories of a selection statement?
10. Between what two language characteristics is a trade-off made when deciding whether more than one selectable segment is executed in one execution of a multiple selection statement?
11. What is the role of the **default** segment in a **switch** statement?
12. What functional languages do not have statements as selectors?
13. Explain how C#’s switch statement is safer than that of C.
14. Under what circumstances can a linear search on a table of cases and labels be accepted?
15. What mechanism does a compiler follow when the number of cases in a selection statement is 10 or greater to optimize the time required to execute?
16. What is a loop variable? What is stepsize?
17. What is the difference between a pretest version and a posttest version of a logical loop?
18. In C, what is the significance of a **for** loop without the second expression?

19. What does the **range** function in Python do?

20. What alternative is provided for goto in Java?

21. What are the design issues for logically controlled loop statements?

22. What is the main reason user-located loop control statements were invented?

23. What predefined iterator for standard data structures is available in Perl?

24. What is the use of a **break** statement in switch statements?

25. What are the differences between the **break** statement of C++ and that of Java?

26. What are fatbars in guarded clauses?

27. What is the limitation of a guarded command?

28. How does a functional language implement repetition?

29. How are iterators implemented in Ruby?

30. What language predefines iterators that can be explicitly called to iterate over its predefined data structures?

31. What common programming language borrows part of its design from Dijkstra’s guarded commands?

**P R O B L E M  S E T**

1. Describe three situations where a combined counting and logical looping statement is needed.

2. Study the iterator feature of CLU in Liskov et al. (1981) and determine its advantages and disadvantages.

3. Compare the set of Ada control statements with those of C# and decide which are better and why.

4. What are the pros and cons of using unique closing reserved words on compound statements?

5. What are the arguments, pros and cons, for Python’s use of indentation to specify compound statements in control statements?

6. Analyze the potential readability problems with using closure reserved words for control statements that are the reverse of the corresponding initial reserved words, such as the **case-esac** reserved words of ALGOL 68. For example, consider common typing errors such as transposing characters.

7. Use the *Science Citation Index* to find an article that refers to Knuth (1974). Read the article and Knuth’s paper and write a paper that summarizes both sides of the goto issue.
8. In his paper on the goto issue, Knuth (1974) suggests a loop control statement that allows multiple exits. Read the paper and write an operational semantics description of the statement.

9. What are the arguments both for and against the exclusive use of Boolean expressions in the control statements in Java (as opposed to also allowing arithmetic expressions, as in C++)?

10. Describe a programming situation in which the else clause in Python’s for statement would be convenient.

11. Describe three specific programming situations that require a posttest loop.

12. Speculate as to the reason control can be transferred into a C loop statement.

**Programming Exercises**

1. Rewrite the following pseudocode segment using a loop structure in the specified languages:

   ```
   k = (j + 13) / 27
   loop:
     if k > 10 then goto out
     k = k + 1
     i = 3 * k - 1
     goto loop
   out: ...
   ```

   a. C, C++, Java, or C#
   b. Python
   c. Ruby

   Assume all variables are integer type. Discuss which language, for this code, has the best writability, the best readability, and the best combination of the two.

2. Redo Programming Exercise 1, except this time make all the variables and constants floating-point type, and change the statement

   ```
   k = k + 1
   ```

to

   ```
   k = k + 1.2
   ```

3. Rewrite the following code segment using a multiple-selection statement in the following languages:

   ```
   if ((k == 1) || (k == 2)) j = 2 * k - 1
   if ((k == 3) || (k == 5)) j = 3 * k + 1
   if (k == 4) j = 4 * k - 1
   if ((k == 6) || (k == 7) || (k == 8)) j = k - 2
   ```
a. C, C++, Java, or C#

b. Python

c. Ruby

Assume all variables are integer type. Discuss the relative merits of the use of these languages for this particular code.

4. Consider the following C program segment. Rewrite it using no gotos or breaks.

```c
j = -3;
for (i = 0; i < 3; i++) {
    switch (j + 2) {
    case 3:
    case 2: j--; break;
    case 0: j += 2; break;
    default: j = 0;
    }
    if (j > 0) break;
    j = 3 - i
}
```

5. In a letter to the editor of CACM, Rubin (1987) uses the following code segment as evidence that the readability of some code with gotos is better than the equivalent code without gotos. This code finds the first row of an \( n \times n \) integer matrix named \( x \) that has nothing but zero values.

```c
for (i = 1; i <= n; i++) {
    for (j = 1; j <= n; j++)
        if (x[i][j] != 0)
            goto reject;
    println ('First all-zero row is:', i);
    break;
reject:
}
```

Rewrite this code without gotos in one of the following languages: C, C++, Java, or C#. Compare the readability of your code to that of the example code.

6. Consider the following programming problem: The values of three integer variables—first, second, and third—must be placed in the three variables max, mid, and min, with the obvious meanings, without using arrays or user-defined or predefined subprograms. Write two solutions to this problem, one that uses nested selections and one that does not. Compare the complexity and expected reliability of the two.

7. Rewrite the C program segment of Programming Exercise 4 using if and goto statements in C.

8. Rewrite the C program segment of Programming Exercise 4 in Java without using a switch statement.
9. Translate the following call to Scheme’s \texttt{COND} to C and set the resulting value to $y$.

\begin{verbatim}
(COND
  ((> x 10) x)
  ((< x 5) (* 2 x))
  ((= x 7) (+ x 10)))
\end{verbatim}
Subprograms

9.1 Introduction
9.2 Fundamentals of Subprograms
9.3 Design Issues for Subprograms
9.4 Local Referencing Environments
9.5 Parameter-Passing Methods
9.6 Parameters That Are Subprograms
9.7 Calling Subprograms Indirectly
9.8 Design Issues for Functions
9.9 Overloaded Subprograms
9.10 Generic Subprograms
9.11 User-Defined Overloaded Operators
9.12 Closures
9.13 Coroutines
Subprograms are the fundamental building blocks of programs and are therefore among the most important concepts in programming language design. We now explore the design of subprograms, including parameter-passing methods, local referencing environments, overloaded subprograms, generic subprograms, and the aliasing and problematic side effects that are associated with subprograms. We also include discussions of indirectly called subprograms, closures, and coroutines.

Implementation methods for subprograms are discussed in Chapter 10.

9.1 Introduction

Two fundamental abstraction facilities can be included in a programming language: process abstraction and data abstraction. In the early history of high-level programming languages, only process abstraction was included. Process abstraction, in the form of subprograms, has been a central concept in all programming languages. In the 1980s, however, many people began to believe that data abstraction was equally important. Data abstraction is discussed in detail in Chapter 11.

The first programmable computer, Babbage’s Analytical Engine, built in the 1840s, had the capability of reusing collections of instruction cards at several different places in a program. In a modern programming language, such a collection of statements is written as a subprogram. This reuse results in savings in memory space and coding time. Such reuse is also an abstraction, for the details of the subprogram’s computation are replaced in a program by a statement that calls the subprogram. Instead of describing how some computation is to be done in a program, that description (the collection of statements in the subprogram) is enacted by a call statement, effectively abstracting away the details. This increases the readability of a program by emphasizing its logical structure while hiding its low-level details.

The methods of object-oriented languages are closely related to the subprograms discussed in this chapter. The primary way methods differ from subprograms is the way they are called and their associations with classes and objects. Although these special characteristics of methods are discussed in Chapter 12, the features they share with subprograms, such as parameters and local variables, are discussed in this chapter.

9.2 Fundamentals of Subprograms

9.2.1 General Subprogram Characteristics

All subprograms discussed in this chapter, except the coroutines described in Section 9.13, have the following characteristics:

- Each subprogram has a single entry point.
• The calling program unit is suspended during the execution of the called subprogram, which implies that there is only one subprogram in execution at any given time.
• Control always returns to the caller when the subprogram execution terminates.

Alternatives to these result in coroutines and concurrent units (Chapter 13).
Most subprograms have names, although some are anonymous. Section 9.12 has examples of anonymous subprograms in C#.

9.2.2 Basic Definitions

A subprogram definition describes the interface to and the actions of the subprogram abstraction. A subprogram call is the explicit request that a specific subprogram be executed. A subprogram is said to be active if, after having been called, it has begun execution but has not yet completed that execution. The two fundamental kinds of subprograms, procedures and functions, are defined and discussed in Section 9.2.4.

A subprogram header, which is the first part of the definition, serves several purposes. First, it specifies that the following syntactic unit is a subprogram definition of some particular kind. In languages that have more than one kind of subprogram, the kind of the subprogram is usually specified with a special word. Second, if the subprogram is not anonymous, the header provides a name for the subprogram. Third, it may specify a list of parameters.

Consider the following header examples:

```
def adder(parameters):
```
This is the header of a Python subprogram named `adder`. Ruby subprogram headers also begin with `def`. The header of a JavaScript subprogram begins with `function`.

In C, the header of a function named `adder` might be as follows:

```
void adder (parameters)
```
The reserved word `void` in this header indicates that the subprogram does not return a value.

The body of subprograms defines its actions. In the C-based languages (and some others—for example, JavaScript) the body of a subprogram is delimited by braces. In Ruby, an `end` statement terminates the body of a subprogram. As with compound statements, the statements in the body of a Python function must be indented and the end of the body is indicated by the first statement that is not indented.

One characteristic of Python functions that sets them apart from the functions of other common programming languages is that function `def` statements are executable. When a `def` statement is executed, it assigns the given name to

1. Some programming languages include both kinds of subprograms, procedures and functions.
the given function body. Until a function’s `def` has been executed, the function cannot be called. Consider the following skeletal example:

```python
if . . .
   def fun(. . .):
      . . .
else
   def fun(. . .):
      . . .
```

If the then clause of this selection construct is executed, that version of the function `fun` can be called, but not the version in the else clause. Likewise, if the else clause is chosen, its version of the function can be called but the one in the then clause cannot.

Ruby methods differ from the subprograms of other programming languages in several interesting ways. Ruby methods are often defined in class definitions but can also be defined outside class definitions, in which case they are considered methods of the root object, `Object`. Such methods can be called without an object receiver, as if they were functions in C or C++. If a Ruby method is called without a receiver, `self` is assumed. If there is no method by that name in the class, enclosing classes are searched, up to `Object`, if necessary.

All Lua functions are anonymous, although they can be defined using syntax that makes it appear as though they have names. For example, consider the following identical definitions of the function `cube`:

```lua
function cube(x) return x * x * x end

cube = function (x) return x * x * x end
```

The first of these uses conventional syntax, while the form of the second more accurately illustrates the namelessness of functions.

The parameter profile of a subprogram contains the number, order, and types of its formal parameters. The protocol of a subprogram is its parameter profile plus, if it is a function, its return type. In languages in which subprograms have types, those types are defined by the subprogram’s protocol.

Subprograms can have declarations as well as definitions. This form parallels the variable declarations and definitions in C, in which declarations are used to provide type information but not to define variables. Subprogram declarations provide the subprogram’s protocol but do not include their bodies. They are necessary in languages that do not allow forward references to subprograms. In both the cases of variables and subprograms, declarations are needed for static type checking. In the case of subprograms, it is the type of the parameters that must be checked. Function declarations are common in C and C++ programs, where they are called prototypes. Such declarations are often placed in header files.
In most other languages (other than C and C++), subprograms do not need declarations, because there is no requirement that subprograms be defined before they are called.

### 9.2.3 Parameters

Subprograms typically describe computations. There are two ways that a nonmethod subprogram can gain access to the data that it is to process: through direct access to nonlocal variables (declared elsewhere but visible in the subprogram) or through parameter passing. Data passed through parameters are accessed using names that are local to the subprogram. Parameter passing is more flexible than direct access to nonlocal variables. In essence, a subprogram with parameter access to the data that it is to process is a parameterized computation. It can perform its computation on whatever data it receives through its parameters (presuming the types of the parameters are as expected by the subprogram). If data access is through nonlocal variables, the only way the computation can proceed on different data is to assign new values to those nonlocal variables between calls to the subprogram. Extensive access to nonlocals can reduce reliability. Variables that are visible to the subprogram where access is desired often end up also being visible where access to them is not needed. This problem was discussed in Chapter 5.

Although methods also access external data through nonlocal references and parameters, the primary data to be processed by a method is the object through which the method is called. However, when a method does access nonlocal data, the reliability problems are the same as with nonmethod subprograms. Also, in an object-oriented language, method access to class variables (those associated with the class, rather than an object) is related to the concept of nonlocal data and should be avoided whenever possible. In this case, as well as the case of a C function accessing nonlocal data, the method can have the side effect of changing something other than its parameters or local data. Such changes complicate the semantics of the method and make it less reliable.

Pure functional programming languages, such as Haskell, do not have mutable data, so functions written in them are unable to change memory in any way—they simply perform calculations and return a resulting value (or function, since functions are values in a pure functional language).

In some situations, it is convenient to be able to transmit computations, rather than data, as parameters to subprograms. In these cases, the name of the subprogram that implements that computation may be used as a parameter. This form of parameter is discussed in Section 9.6. Data parameters are discussed in Section 9.5.

The parameters in the subprogram header are called **formal parameters**. They are sometimes thought of as dummy variables because they are not variables in the usual sense: In most cases, they are bound to storage only when the subprogram is called, and that binding is often through some other program variables.
Subprogram call statements must include the name of the subprogram and a list of parameters to be bound to the formal parameters of the subprogram. These parameters are called actual parameters. They must be distinguished from formal parameters, because the two usually have different restrictions on their forms, and of course, their uses are quite different.

In most programming languages, the correspondence between actual and formal parameters—or the binding of actual parameters to formal parameters—is done by position: The first actual parameter is bound to the first formal parameter and so forth. Such parameters are called positional parameters. This is an effective and safe method of relating actual parameters to their corresponding formal parameters, as long as the parameter lists are relatively short.

When parameter lists are long, however, it is easy for a programmer to make mistakes in the order of actual parameters in the list. One solution to this problem is to provide keyword parameters, in which the name of the formal parameter to which an actual parameter is to be bound is specified with the actual parameter in a call. The advantage of keyword parameters is that they can appear in any order in the actual parameter list. Python functions can be called using this technique, as in

```python
sumer(length = my_length,
       list = my_array,
       sum = my_sum)
```

where the definition of `sumer` has the formal parameters `length`, `list`, and `sum`.

The disadvantage to keyword parameters is that the user of the subprogram must know the names of formal parameters.

In addition to keyword parameters, some languages, for example Python, allow positional parameters. Keyword and positional parameters can be mixed in a call, as in

```python
sumer(my_length,
      sum = my_sum,
      list = my_array)
```

The only restriction with this approach is that after a keyword parameter appears in the list, all remaining parameters must be keyworded. This restriction is necessary because a position may no longer be well defined after a keyword parameter has appeared.

In Python, Ruby, C++, and PHP, formal parameters can have default values. A default value is used if no actual parameter is passed to the formal parameter in the subprogram header. Consider the following Python function header:

```python
def compute_pay(income, exemptions = 1, tax_rate)
```

2. Some authors call actual parameters arguments and formal parameters just parameters.
The exemptions formal parameter can be absent in a call to `compute_pay`; when it is, the value 1 is used. No comma is included for an absent actual parameter in a Python call, because the only value of such a comma would be to indicate the position of the next parameter, which in this case is not necessary because all actual parameters after an absent actual parameter must be keyworded. For example, consider the following call:

```python
pay = compute_pay(20000.0, tax_rate = 0.15)
```

In C++, which does not support keyword parameters, the rules for default parameters are necessarily different. The default parameters must appear last, because parameters are positionally associated. Once a default parameter is omitted in a call, all remaining formal parameters must have default values. A C++ function header for the `compute_pay` function can be written as follows:

```cpp
float compute_pay(float income, float tax_rate,
                  int exemptions = 1)
```

Notice that the parameters are rearranged so that the one with the default value is last. An example call to the C++ `compute_pay` function is

```cpp
pay = compute_pay(20000.0, 0.15);
```

In most languages that do not have default values for formal parameters, the number of actual parameters in a call must match the number of formal parameters in the subprogram definition header. However, in C, C++, Perl, JavaScript, and Lua this is not required. When there are fewer actual parameters in a call than formal parameters in a function definition, it is the programmer's responsibility to ensure that the parameter correspondence, which is always positional, and the subprogram execution are sensible.

Although this design, which allows a variable number of parameters, is clearly prone to error, it is also sometimes convenient. For example, the `printf` function of C can print any number of items (data values and/or literals).

C# allows methods to accept a variable number of parameters, as long as they are of the same type. The method specifies its formal parameter with the `params` modifier. The call can send either an array or a list of expressions, whose values are placed in an array by the compiler and provided to the called method. For example, consider the following method:

```csharp
public void DisplayList(params int[] list) {
    foreach (int next in list) {
        Console.WriteLine("Next value {0}", next);
    }
}
```
If `DisplayList` is defined for the class `MyClass` and we have the following declarations,

```ruby
Myclass myObject = new MyClass;
int[] myList = new int[6] {2, 4, 6, 8, 10, 12};
```

`DisplayList` could be called with either of the following:

```ruby
myObject.DisplayList(myList);
myObject.DisplayList(2, 4, 3 * x - 1, 17);
```

Ruby supports a complicated but highly flexible actual parameter configuration. The initial parameters are expressions, whose value objects are passed to the corresponding formal parameters. The initial parameters can be followed by a list of key => value pairs, which are placed in an anonymous hash and a reference to that hash is passed to the next formal parameter. These are used as a substitute for keyword parameters, which Ruby does not support. The hash item can be followed by a single parameter preceded by an asterisk. This parameter is called the `array formal parameter`. When the method is called, the array formal parameter is set to reference a new `Array` object. All remaining actual parameters are assigned to the elements of the new `Array` object. If the actual parameter that corresponds to the array formal parameter is an array, it must also be preceded by an asterisk, and it must be the last actual parameter.³

So, Ruby allows a variable number of parameters in a way similar to that of C#. Because Ruby arrays can store different types, there is no requirement that the actual parameters passed to the array have the same type.

The following example skeletal function definition and call illustrate the parameter structure of Ruby:

```ruby
list = [2, 4, 6, 8]
def tester(p1, p2, p3, *p4)
    ...
end ...
tester('first', mon => 72, tue => 68, wed => 59, *list)
```

Inside `tester`, the values of its formal parameters are as follows:

- `p1` is `'first'`
- `p2` is `{mon => 72, tue => 68, wed => 59}`
- `p3` is `2`
- `p4` is `[4, 6, 8]`

Python supports parameters that are similar to those of Ruby.

³. Not quite true, because the array formal parameter can be followed by a method or function reference, which is preceded by an ampersand (&).
Lua uses a simple mechanism for supporting a variable number of parameters—such parameters are represented by an ellipsis (...). This ellipsis can be treated as an array or as a list of values that can be assigned to a list of variables. For example, consider the following two function examples:

```lua
function multiply (. . .)
    local product = 1
    for i, next in ipairs{. . .} do
        product = product * next
    end
    return sum
end
```

`ipairs` is an iterator for arrays (it returns the index and value of the elements of an array, one element at a time). {...} is an array of the actual parameter values.

```lua
function DoIt (. . .)
    local a, b, c = . . .
    . . .
end
```

Suppose `DoIt` is called with the following call:

doit(4, 7, 3)

In this example, `a`, `b`, and `c` will be initialized in the function to the values 4, 7, and 3, respectively.

The three-period parameter need not be the only parameter—it can appear at the end of a list of named formal parameters.

### 9.2.4 Procedures and Functions

There are two distinct categories of subprograms—procedures and functions—both of which can be viewed as approaches to extending the language. Subprograms are collections of statements that define parameterized computations. Functions return values and procedures do not. In most languages that do not include procedures as a separate form of subprogram, functions can be defined not to return values and they can be used as procedures. The computations of a procedure are enacted by single call statements. In effect, procedures define new statements. For example, if a particular language does not have a sort statement, a user can build a procedure to sort arrays of data and use a call to that procedure in place of the unavailable sort statement. Only some older languages, such as Fortran and Ada, support procedures.

Procedures can produce results in the calling program unit by two methods: (1) If there are variables that are not formal parameters but are still visible in both the procedure and the calling program unit, the procedure can change them; and (2) if the procedure has formal parameters that allow the transfer of data to the caller, those parameters can be changed.
Functions structurally resemble procedures but are semantically modeled on mathematical functions. If a function is a faithful model, it produces no side effects; that is, it modifies neither its parameters nor any variables defined outside the function. Such a function returns a value—that is its only desired effect. The functions in most programming languages have side effects.

Functions are called by appearances of their names in expressions, along with the required actual parameters. The value produced by a function’s execution is returned to the calling code, effectively replacing the call itself. For example, the value of the expression \( f(x) \) is whatever value \( f \) produces when called with the parameter \( x \). For a function that does not produce side effects, the returned value is its only effect.

Functions define new user-defined operators. For example, if a language does not have an exponentiation operator, a function can be written that returns the value of one of its parameters raised to the power of another parameter. Its header in C++ could be

```cpp
float power(float base, float exp)
```

which could be called with

```cpp
result = 3.4 * power(10.0, x)
```

The standard C++ library already includes a similar function named `pow`. Compare this with the same operation in Perl, in which exponentiation is a built-in operation:

```cpp
result = 3.4 * 10.0 ** x
```

In some programming languages, users are permitted to overload operators by defining new functions for operators. User-defined overloaded operators are discussed in Section 9.11.

## 9.3 Design Issues for Subprograms

Subprograms are complex structures, and it follows from this that a lengthy list of issues is involved in their design. One obvious issue is the choice of one or more parameter-passing methods that will be used. The wide variety of approaches that have been used in various languages is a reflection of the diversity of opinion on the subject. A closely related issue is whether the types of actual parameters will be type checked against the types of the corresponding formal parameters.

The nature of the local environment of a subprogram dictates to some degree the nature of the subprogram. The most important question here is whether local variables are statically or dynamically allocated.

Next, there is the question of whether subprogram definitions can be nested. Another issue is whether subprogram names can be passed as parameters. If
subprogram names can be passed as parameters and the language allows sub-
programs to be nested, there is the question of the correct referencing envi-
ronment of a subprogram that has been passed as a parameter.

As seen in Chapter 5, side effects of functions can cause problems. So, restrictions on side effects are a design issue for functions. The types and num-
ber of values that can be returned from functions are other design issues.

Finally, there are the questions of whether subprograms can be overloaded or generic. An **overloaded subprogram** is one that has the same name as another subprogram in the same referencing environment. A **generic subpro-
gram** is one whose computation can be done on data of different types in different calls. A **closure** is a nested subprogram and its referencing environment, which together allow the subprogram to be called from anywhere in a program.

The following is a summary of these design issues for subprograms in general. Additional issues that are specifically associated with functions are discussed in Section 9.10.

- Are local variables statically or dynamically allocated?
- Can subprogram definitions appear in other subprogram definitions?
- What parameter-passing method or methods are used?
- Are the types of the actual parameters checked against the types of the formal parameters?
- If subprograms can be passed as parameters and subprograms can be nested, what is the referencing environment of a passed subprogram?
- Are functional side effects allowed?
- What types of values can be returned from functions?
- How many values can be returned from functions?
- Can subprograms be overloaded?
- Can subprograms be generic?
- If the language allows nested subprograms, are closures supported?

These issues and example designs are discussed in the following sections.

### 9.4 Local Referencing Environments

This section discusses the issues related to variables that are defined within sub-
programs. The issue of nested subprogram definitions is also briefly covered.

#### 9.4.1 Local Variables

Subprograms can define their own variables, thereby defining local referencing environments. Variables that are defined inside subprograms are called **local variables**, because their scope is usually the body of the subprogram in which they are defined.
In the terminology of Chapter 5, local variables can be either static or stack dynamic. If local variables are stack dynamic, they are bound to storage when the subprogram begins execution and are unbound from storage when that execution terminates. There are several advantages of stack-dynamic local variables, the primary one being flexibility. It is essential that recursive subprograms have stack-dynamic local variables. Another advantage of stack-dynamic locals is that the storage for local variables in an active subprogram can be shared with the local variables in all inactive subprograms. This is not as important an advantage as it was when computers had smaller memories.

The main disadvantages of stack-dynamic local variables are the following: First, there is the cost of the time required to allocate, initialize (when necessary), and deallocate such variables for each call to the subprogram. Second, accesses to stack-dynamic local variables must be indirect, whereas accesses to static variables can be direct. This indirectness is required because the place in the stack where a particular local variable will reside can be determined only during execution (see Chapter 10). Finally, when all local variables are stack dynamic, subprograms cannot be history sensitive; that is, they cannot retain data values of local variables between calls. It is sometimes convenient to be able to write history-sensitive subprograms. A common example of a need for a history-sensitive subprogram is one whose task is to generate pseudorandom numbers. Each call to such a subprogram computes one pseudorandom number, using the last one it computed. It must, therefore, store the last one in a static local variable. Coroutines and the subprograms used in iterator loop constructs (discussed in Chapter 8) are other examples of subprograms that need to be history sensitive.

The primary advantage of static local variables over stack-dynamic local variables is that they are slightly more efficient—they require no run-time overhead for allocation and deallocation. Also, if accessed directly, these accesses are obviously more efficient. And, of course, they allow subprograms to be history sensitive. The greatest disadvantage of static local variables is their inability to support recursion. Also, their storage cannot be shared with the local variables of other inactive subprograms.

In most contemporary languages, local variables in a subprogram are by default stack dynamic. In C and C++ functions, locals are stack dynamic unless specifically declared to be static. For example, in the following C (or C++) function, the variable sum is static and count is stack dynamic.

```c
int adder(int list[], int listlen) {
    static int sum = 0;
    int count;
    for (count = 0; count < listlen; count ++)
        sum += list [count];
    return sum;
}
```

4. In some implementations, static variables are also accessed indirectly, thereby eliminating this disadvantage.
The methods of C++, Java, and C# have only stack-dynamic local variables. In Python, the only declarations used in method definitions are for globals. Any variable declared to be global in a method must be a variable defined outside the method. A variable defined outside the method can be referenced in the method without declaring it to be global, but such a variable cannot be assigned in the method. If the name of a global variable is assigned in a method, it is implicitly declared to be a local and the assignment does not disturb the global. All local variables in Python methods are stack dynamic.

Only variables with restricted scope are declared in Lua. Any block, including the body of a function, can declare local variables with the `local` declaration, as in the following:

```
local sum
```

All nondeclared variables in Lua are global. Access to local variables in Lua is faster than access to global variables according to Ierusalimschy (2006).

### 9.4.2 Nested Subprograms

The idea of nesting subprograms originated with ALGOL 60. The motivation was to be able to create a hierarchy of both logic and scopes. If a subprogram is needed only within another subprogram, why not place it there and hide it from the rest of the program? Because static scoping is usually used in languages that allow subprograms to be nested, this also provides a highly structured way to grant access to nonlocal variables in enclosing subprograms. Recall that in Chapter 5, the problems introduced by this were discussed. For a long time, the only languages that allowed nested subprograms were those directly descending from ALGOL 60, which were ALGOL 68, Pascal, and Ada. Many other languages, including all of the direct descendants of C, do not allow subprogram nesting. Recently, some new languages again allow it. Among these are JavaScript, Python, Ruby, and Lua. Also, most functional programming languages allow subprograms to be nested.

### 9.5 Parameter-Passing Methods

Parameter-passing methods are the ways in which parameters are transmitted to and/or from called subprograms. First, we focus on the different semantics models of parameter-passing methods. Then, we discuss the various implementation models invented by language designers for these semantics models. Next, we survey the design choices of several languages and discuss the actual methods used to implement the implementation models. Finally, we consider the design considerations that face a language designer in choosing among the methods.
9.5.1 Semantics Models of Parameter Passing

Formal parameters are characterized by one of three distinct semantics models: (1) They can receive data from the corresponding actual parameter; (2) they can transmit data to the actual parameter; or (3) they can do both. These models are called **in mode**, **out mode**, and **inout mode**, respectively. For example, consider a subprogram that takes two arrays of int values as parameters—list1 and list2. The subprogram must add list1 to list2 and return the result as a revised version of list2. Furthermore, the subprogram must create a new array from the two given arrays and return it. For this subprogram, list1 should be in mode, because it is not to be changed by the subprogram. list2 must be inout mode, because the subprogram needs the given value of the array and must return its new value. The third array should be out mode, because there is no initial value for this array and its computed value must be returned to the caller.

There are two conceptual models of how data transfers take place in parameter transmission: Either an actual value is copied (to the caller, to the called, or both ways) or an access path is transmitted. Most commonly, the access path is a simple pointer or reference. Figure 9.1 illustrates the three semantics models of parameter passing when values are copied.

9.5.2 Implementation Models of Parameter Passing

A variety of models have been developed by language designers to guide the implementation of the three basic parameter transmission modes. In the following sections, we discuss several of these, along with their relative strengths and weaknesses.

---

**Figure 9.1**

The three semantics models of parameter passing when physical moves are used.
9.5.2.1 Pass-by-Value

When a parameter is passed by value, the value of the actual parameter is used to initialize the corresponding formal parameter, which then acts as a local variable in the subprogram, thus implementing in-mode semantics.

Pass-by-value is normally implemented by copy, because accesses often are more efficient with this approach. It could be implemented by transmitting an access path to the value of the actual parameter in the caller, but that would require that the value be in a write-protected cell (one that can only be read). Enforcing the write protection is not always a simple matter. For example, suppose the subprogram to which the parameter was passed passes it in turn to another subprogram. This is another reason to use copy transfer. As we will see in Section 9.5.4, C++ provides a convenient and effective method for specifying write protection on pass-by-value parameters that are transmitted by access path.

The advantage of pass-by-value is that for scalars it is fast, in both linkage cost and access time.

The main disadvantage of the pass-by-value method if copies are used is that additional storage is required for the formal parameter, either in the called subprogram or in some area outside both the caller and the called subprogram. In addition, the actual parameter must be copied to the storage area for the corresponding formal parameter. The storage and the copy operations can be costly if the parameter is large, such as an array with many elements.

9.5.2.2 Pass-by-Result

Pass-by-result is an implementation model for out-mode parameters. When a parameter is passed by result, no value is transmitted to the subprogram. The corresponding formal parameter acts as a local variable, but just before control is transferred back to the caller, its value is transmitted back to the caller's actual parameter, which obviously must be a variable. (How would the caller reference the computed result if it were a literal or an expression?)

The pass-by-result method has the advantages and disadvantages of pass-by-value, plus some additional disadvantages. If values are returned by copy (as opposed to access paths), as they typically are, pass-by-result also requires the extra storage and the copy operations that are required by pass-by-value. As with pass-by-value, the difficulty of implementing pass-by-result by transmitting an access path usually results in it being implemented by copy. In this case, the problem is in ensuring that the initial value of the actual parameter is not used in the called subprogram.

One additional problem with the pass-by-result model is that there can be an actual parameter collision, such as the one created with the call

\[ \text{sub(p1, p1)} \]

In sub, assuming the two formal parameters have different names, the two can obviously be assigned different values. Then, whichever of the two is copied
to their corresponding actual parameter last becomes the value of \( p1 \) in the caller. Thus, the order in which the actual parameters are copied determines their value. For example, consider the following C# method, which specifies the pass-by-result method with the \texttt{out} specifier on its formal parameter:\footnote{The \texttt{out} specifier must also be specified on the corresponding actual parameter.}

```csharp
void Fixer(out int x, out int y) {
    x = 17;
    y = 35;
}

f.Fixer(out a, out a);
```

If, at the end of the execution of \texttt{Fixer}, the formal parameter \( x \) is assigned to its corresponding actual parameter first, then the value of the actual parameter \( a \) in the caller will be 35. If \( y \) is assigned first, then the value of the actual parameter \( a \) in the caller will be 17.

Because the order can be implementation dependent for some languages, different implementations can produce different results.

Calling a procedure with two identical actual parameters can also lead to different kinds of problems when other parameter-passing methods are used, as discussed in Section 9.5.2.4.

Another problem that can occur with pass-by-result is that the implementor may be able to choose between two different times to evaluate the addresses of the actual parameters: at the time of the call or at the time of the return. For example, consider the following C# method and following code:

```csharp
void DoIt(out int x, int index){
    x = 17;
    index = 42;
}

sub = 21;
f.DoIt(list[sub], sub);
```

The address of \texttt{list[sub]} changes between the beginning and end of the method. The implementor must choose the time to bind this parameter to an address—at the time of the call or at the time of the return. If the address is computed on entry to the method, the value 17 will be returned to \texttt{list[21]}; if computed just before return, 17 will be returned to \texttt{list[42]}. This makes programs unportable between an implementation that chooses to evaluate the addresses for out-mode parameters at the beginning of a subprogram and one that chooses to do that evaluation at the end. An obvious way to avoid this problem is for the language designer to specify when the address to be used to return the parameter value must be computed.
9.5.2.3 Pass-by-Value-Result

Pass-by-value-result is an implementation model for inout-mode parameters in which actual values are copied. It is in effect a combination of pass-by-value and pass-by-result. The value of the actual parameter is used to initialize the corresponding formal parameter, which then acts as a local variable. In fact, pass-by-value-result formal parameters must have local storage associated with the called subprogram. At subprogram termination, the value of the formal parameter is transmitted back to the actual parameter.

Pass-by-value-result is sometimes called pass-by-copy, because the actual parameter is copied to the formal parameter at subprogram entry and then copied back at subprogram termination.

Pass-by-value-result shares with pass-by-value and pass-by-result the disadvantages of requiring multiple storage for parameters and time for copying values. It shares with pass-by-result the problems associated with the order in which actual parameters are assigned.

The advantages of pass-by-value-result are relative to pass-by-reference, so they are discussed in Section 9.5.2.4.

9.5.2.4 Pass-by-Reference

Pass-by-reference is a second implementation model for inout-mode parameters. Rather than copying data values back and forth, however, as in pass-by-value-result, the pass-by-reference method transmits an access path, usually just an address, to the called subprogram. This provides the access path to the cell storing the actual parameter. Thus, the called subprogram is allowed to access the actual parameter in the calling program unit. In effect, the actual parameter is shared with the called subprogram.

The advantage of pass-by-reference is that the passing process itself is efficient, in terms of both time and space. Duplicate space is not required and no copying is required.

There are, however, several disadvantages to the pass-by-reference method. First, access to the formal parameters will be slower than pass-by-value parameters, because of the additional level of indirect addressing that is required. Second, if only one-way communication to the called subprogram is required, inadvertent and erroneous changes may be made to the actual parameter.

Another problem of pass-by-reference is that aliases can be created. This problem should be expected, because pass-by-reference makes access paths available to the called subprograms, thereby providing access to nonlocal variables. The problem with these kinds of aliasing is the same as in other circumstances: It is harmful to readability and thus to reliability. It also makes program verification more difficult.

There are several ways pass-by-reference parameters can create aliases. First, collisions can occur between actual parameters. Consider a C++ function that has two parameters that are to be passed by reference, as in

---

6. This is further explained in Section 9.5.3.
void fun(int &first, int &second)

If the call to fun happens to pass the same variable twice, as in

fun(total, total)

then first and second in fun will be aliases.

Second, collisions between array elements can also cause aliases. For example, suppose the function fun is called with two array elements that are specified with variable subscripts, as in

fun(list[i], list[j])

If these two parameters are passed by reference and i happens to be equal to j, then first and second are again aliases.

Third, if two of the formal parameters of a subprogram are an element of an array and the whole array, and both are passed by reference, then a call such as

fun1(list[i], list)

could result in aliasing in fun1, because fun1 can access all elements of list through the second parameter and access a single element through its first parameter.

Still another way to get aliasing with pass-by-reference parameters is through collisions between formal parameters and nonlocal variables that are visible. For example, consider the following C code:

```c
int * global;
void main() {
    ... 
    sub(global);
    ... }

void sub(int * param) {
    ... }
```

Inside sub, param and global are aliases.

All these possible aliasing situations are eliminated if pass-by-value-result is used instead of pass-by-reference. However, in place of aliasing, other problems sometimes arise, as discussed in Section 9.5.2.3.

### 9.5.2.5 Pass-by-Name

**Pass-by-name** is an inout-mode parameter transmission method that does not correspond to a single implementation model. When parameters are passed by name, the actual parameter is, in effect, textually substituted for the corresponding
formal parameter in all its occurrences in the subprogram. This method is quite different from those discussed thus far; in which case, formal parameters are bound to actual values or addresses at the time of the subprogram call. A pass-by-name formal parameter is bound to an access method at the time of the subprogram call, but the actual binding to a value or an address is delayed until the formal parameter is assigned or referenced. Implementing a pass-by-name parameter requires a subprogram to be passed to the called subprogram to evaluate the address or value of the formal parameter. The referencing environment of the passed subprogram must also be passed. This subprogram/referencing environment is a closure (see Section 9.12). Pass-by-name parameters are both complex to implement and inefficient. They also add significant complexity to the program, thereby lowering its readability and reliability.

Because pass-by-name is not part of any widely used language, it is not discussed further here. However, it is used at compile time by the macros in assembly languages and for the generic parameters of the generic subprograms in C++, Java 5.0, and C# 2005, as discussed in Section 9.9.

9.5.3 Implementing Parameter-Passing Methods

We now address the question of how the various implementation models of parameter passing are actually implemented.

In most contemporary languages, parameter communication takes place through the run-time stack. The run-time stack is initialized and maintained by the run-time system, which manages the execution of programs. The run-time stack is used extensively for subprogram control linkage and parameter passing, as discussed in Chapter 10. In the following discussion, we assume that the stack is used for all parameter transmission.

Pass-by-value parameters have their values copied into stack locations. The stack locations then serve as storage for the corresponding formal parameters. Pass-by-result parameters are implemented as the opposite of pass-by-value. The values assigned to the pass-by-result actual parameters are placed in the stack, where they can be retrieved by the calling program unit upon termination of the called subprogram. Pass-by-value-result parameters can be implemented directly from their semantics as a combination of pass-by-value and pass-by-result. The stack location for such a parameter is initialized by the call and is then used like a local variable in the called subprogram.

Pass-by-reference parameters are perhaps the simplest to implement. Regardless of the type of the actual parameter, only its address must be placed in the stack. In the case of literals, the address of the literal is put in the stack. In the case of an expression, the compiler must build code to evaluate the expression, which must be executed just before the transfer of control to the called subprogram. The address of the memory cell in which the code places the result of its evaluation is then put in the stack. The compiler must be sure to prevent the called subprogram from changing parameters that are literals or expressions.

7. These closures were originally (in ALGOL 60) called *thunks*. 
Figure 9.2

One possible stack implementation of the common parameter-passing methods

Function header: **void** sub (int a, int b, int c, int d)
Function call in main: sub (w, x, y, z)
(pass w by value, x by result, y by value-result, z by reference)

Access to the formal parameters in the called subprogram is by indirect addressing from the stack location of the address. The implementation of pass-by-value, -result, -value-result, and -reference, where the run-time stack is used, is shown in Figure 9.2. Subprogram sub is called from main with the call sub (w, x, y, z), where w is passed by value, x is passed by result, y is passed by value-result, and z is passed by reference.

### 9.5.4 Parameter-Passing Methods of Some Common Languages

C uses pass-by-value. Pass-by-reference (inout mode) semantics is achieved by using pointers as parameters. The value of the pointer is made available to the called function and nothing is copied back. However, because what was passed is an access path to the data of the caller, the called function can change the caller’s data. C copied this use of the pass-by-value method from ALGOL 68. In both C and C++, formal parameters can be typed as pointers to constants. The corresponding actual parameters need not be constants, for in such cases they are coerced to constants. This allows pointer parameters to provide the efficiency of pass-by-reference with the one-way semantics of pass-by-value. Write protection of those parameters in the called function is implicitly specified.

C++ includes a special pointer type, called a *reference type*, as discussed in Chapter 6, which is often used for parameters. Reference parameters are implicitly dereferenced in the function or method, and their semantics is pass-by-reference. C++ also allows reference parameters to be defined to be constants. For example, we could have
void fun(int &p1, int p2, &p3) { ... } where p1 is pass-by-reference but cannot be changed in the function fun, p2 is pass-by-value, and p3 is pass-by-reference. Neither p1 nor p3 need be explicitly dereferenced in fun.

Constant parameters and in-mode parameters are not exactly alike. Constant parameters clearly implement in mode. However, in all of the common imperative languages except Ada, in-mode parameters can be assigned in the subprogram even though those changes are never reflected in the values of the corresponding actual parameters. Constant parameters can never be assigned.

As with C and C++, all Java parameters are passed by value. However, because objects can be accessed only through reference variables, object parameters are in effect passed by reference. Although an object reference passed as a parameter cannot itself be changed in the called subprogram, the referenced object can be changed if a method is available to cause the change. Because reference variables cannot point to scalar variables directly and Java does not have pointers, scalars cannot be passed by reference in Java (although a reference to an object that contains a scalar can).

The default parameter-passing method of C# is pass-by-value. Pass-by-reference can be specified by preceding both a formal parameter and its corresponding actual parameter with ref. For example, consider the following C# skeletal method and call:

```csharp
void sumer(ref int oldSum, int newOne) { ... }
sumer(ref sum, newValue);
```

The first parameter to sumer is passed by reference; the second is passed by value.

C# also supports out-mode parameters, which are pass-by-reference parameters that do not need initial values. Such parameters are specified in the formal parameter list with the out modifier.

PHP’s parameter passing is similar to that of C#, except that either the actual parameter or the formal parameter can specify pass-by-reference. Pass-by-reference is specified by preceding one or both of the parameters with an ampersand.

Perl employs a primitive means of passing parameters. All actual parameters are implicitly placed in a predefined array named @_ (of all things!). The subprogram retrieves the actual parameter values (or addresses) from this array. The most peculiar thing about this array is its magical nature, exposed by the fact that its elements are in effect aliases for the actual parameters. Therefore, if an element of @_ is changed in the called subprogram, that change is reflected in the corresponding actual parameter in the call, assuming there is a corresponding actual parameter (the number of actual parameters need not be the same as the number of formal parameters) and it is a variable.
The parameter-passing method of Python and Ruby is called **pass-by-assignment**. Because all data values are objects, every variable is a reference to an object. In pass-by-assignment, the actual parameter value is assigned to the formal parameter. Therefore, pass-by-assignment is in effect pass-by-reference, because the value of all actual parameters are references. However, only in certain cases does this result in pass-by-reference semantics. For example, many objects are essentially immutable. In a pure object-oriented language, the process of changing the value of a variable with an assignment statement, as in

\[
x = x + 1
\]

does not change the object referenced by \(x\). Rather, it takes the object referenced by \(x\), increments it by 1, thereby creating a new object (with the value \(x + 1\)), and then changes \(x\) to reference the new object. So, when a reference to a scalar object is passed to a subprogram, the object being referenced cannot be changed in place. Because the reference is passed by value, even though the formal parameter is changed in the subprogram, that change has no effect on the actual parameter in the caller.

Now, suppose a reference to an array is passed as a parameter. If the corresponding formal parameter is assigned a new array object, there is no effect on the caller. However, if the formal parameter is used to assign a value to an element of the array, as in

\[
\text{list}[3] = 47
\]

the actual parameter is affected. So, changing the reference of the formal parameter has no effect on the caller, but changing an element of the array that is passed as a parameter does.

### 9.5.5 Type Checking Parameters

It is now widely accepted that software reliability demands that the types of actual parameters be checked for consistency with the types of the corresponding formal parameters. Without such type checking, small typographical errors can lead to program errors that may be difficult to diagnose because they are not detected by the compiler or the run-time system. For example, in the function call

\[
\text{result} = \text{sub1}(1)
\]

the actual parameter is an integer constant. If the formal parameter of \text{sub1} is a floating-point type, no error will be detected without parameter type checking. Although an integer 1 and a floating-point 1 have the same value, the representations of these two are very different. \text{sub1} cannot produce a correct result given an integer actual parameter value if it expects a floating-point value.
Early programming languages, such as Fortran 77 and the original version of C, did not require parameter type checking; most later languages require it. However, the relatively recent languages Perl, JavaScript, and PHP do not.

C and C++ require some special discussion in the matter of parameter type checking. In the original C, neither the number of parameters nor their types were checked. In C89, the formal parameters of functions can be defined in two ways. They can be defined as in the original C; that is, the names of the parameters are listed in parentheses and the type declarations for them follow, as in the following function:

```c
double sin(x)
  double x;
  { . . . }
```

Using this form avoids type checking, thereby allowing calls such as

```c
double value;
int count;
  . . .
value = sin(count);
```

to be legal, although they are never correct.

The alternative to the original C definition approach is called the **prototype** method, in which the formal parameter types are included in the list, as in

```c
double sin(double x)
  { . . . }
```

If this version of `sin` is called with the same call, that is, with the following, it is also legal:

```c
value = sin(count);
```

The type of the actual parameter (`int`) is checked against that of the formal parameter (`double`). Although they do not match, `int` is coercible to `double` (it is a widening coercion), so the conversion is done. If the conversion is not possible (for example, if the actual parameter had been an array) or if the number of parameters is wrong, then a semantics error is detected. So in C89, the user chooses whether parameters are to be type checked.

In C99 and C++, all functions must have their formal parameters in prototype form. However, type checking can be avoided for some of the parameters by replacing the last part of the parameter list with an ellipsis, as in

```c
int printf(const char* format_string, . . .);
```

A call to `printf` must include at least one parameter, a pointer to a literal character string. Beyond that, anything (including nothing) is legal. The way `printf` determines whether there are additional parameters is by the presence
of format codes in the string parameter. For example, the format code for integer output is `%d`. This appears as part of the string, as in the following:

```c
printf("The sum is %d\n", sum);
```

The `%` tells the `printf` function that there is one more parameter.

There is one more interesting issue with actual to formal parameter coercions when primitives can be passed by reference, as in C#. Suppose a call to a method passes a `float` value to a `double` formal parameter. If this parameter is passed by value, the `float` value is coerced to `double` and there is no problem. This particular coercion is very useful, for it allows a library to provide double versions of subprograms that can be used for both `float` and `double` values. However, suppose the parameter is passed by reference. When the value of the `double` formal parameter is returned to the `float` actual parameter in the caller, the value will overflow its location. To avoid this problem, C# requires the type of a `ref` actual parameter to match exactly the type of its corresponding formal parameter (no coercion is allowed).

In Python and Ruby, there is no type checking of parameters, because typing in these languages is a different concept. Objects have types, but variables do not, so formal parameters are typeless. This disallows the very idea of type checking parameters.

### 9.5.6 Multidimensional Arrays as Parameters

The storage-mapping functions that are used to map the index values of references to elements of multidimensional arrays to addresses in memory were discussed at length in Chapter 6. In some languages, such as C and C++, when a multidimensional array is passed as a parameter to a subprogram, the compiler must be able to build the mapping function for that array while seeing only the text of the subprogram (not the calling subprogram). This is true because the subprograms can be compiled separately from the programs that call them. Consider the problem of passing a matrix to a function in C. Multidimensional arrays in C are really arrays of arrays, and they are stored in row major order. Following is a storage-mapping function for row major order for matrices when the lower bound of all indices is 0 and the element size is 1:

```c
address(mat[i, j]) = address(mat[0, 0]) + i *
number_of_columns + j
```

Notice that this mapping function needs the number of columns but not the number of rows. Therefore, in C and C++, when a matrix is passed as a parameter, the formal parameter must include the number of columns in the second pair of brackets. This is illustrated in the following skeletal C program:

```c
void fun(int matrix[][10]) {
  . . .
}
void main() {
```
The problem with this method of passing matrices as parameters is that it does not allow a programmer to write a function that can accept matrices with different numbers of columns; a new function must be written for every matrix with a different number of columns. This, in effect, disallows writing flexible functions that may be effectively reusable if the functions deal with multidimensional arrays. In C and C++, there is a way around the problem because of their inclusion of pointer arithmetic. The matrix can be passed as a pointer, and the actual dimensions of the matrix also can be passed as parameters. Then, the function can evaluate the user-written storage-mapping function using pointer arithmetic each time an element of the matrix must be referenced. For example, consider the following function prototype:

```c
void fun(float *mat_ptr,
        int num_rows,
        int num_cols);
```

The following statement can be used to move the value of the variable \( x \) to the \([row][col]\) element of the parameter matrix in `fun`:

\[
*(\text{mat}\_\text{ptr} + (\text{row} \times \text{num}\_\text{cols} + \text{col})) = x;
\]

Although this works, it is obviously difficult to read, and because of its complexity, it is error prone. The difficulty with reading this can be alleviated by using a macro to define the storage-mapping function, such as

```c
#define mat_ptr(r,c) (*mat_ptr + ((r) \times (num_cols) + (c)))
```

With this, the assignment can be written as

\[
\text{mat}\_\text{ptr}(\text{row},\text{col}) = x;
\]

Other languages use different approaches to dealing with the problem of passing multidimensional arrays.

In Java and C#, arrays are objects. They are all single dimensioned, but the elements can be arrays. Each array inherits a named constant (`length` in Java and `Length` in C#) that is set to the length of the array when the array object is created. The formal parameter for a matrix appears with two sets of empty brackets, as in the following Java method:

```java
float sumer(float mat[][])) {
    float sum = 0.0f;
    
    int mat[5][10];
    . . .
    fun(mat);
    . . .
}
```
Because each array has its own length value, in a matrix the rows can have different lengths.

9.5.7 Design Considerations

Two important considerations are involved in choosing parameter-passing methods: efficiency and whether one-way or two-way data transfer is needed.

Contemporary software-engineering principles dictate that access by subprogram code to data outside the subprogram should be minimized. With this goal in mind, in-mode parameters should be used whenever no data are to be returned through parameters to the caller. Out-mode parameters should be used when no data are transferred to the called subprogram but the subprogram must transmit data back to the caller. Finally, inout-mode parameters should be used only when data must move in both directions between the caller and the called subprogram.

There is a practical consideration that is in conflict with this principle. Sometimes it is justifiable to pass access paths for one-way parameter transmission. For example, when a large array is to be passed to a subprogram that does not modify it, a one-way method may be preferred. However, pass-by-value would require that the entire array be moved to a local storage area of the subprogram. This would be costly in both time and space. Because of this, large arrays are often passed by reference. This is precisely the reason why the Ada 83 definition allowed implementors to choose between the two methods for structured parameters. C++ constant reference parameters offer another solution. Another alternative approach would be to allow the user to choose between the methods.

The choice of a parameter-passing method for functions is related to another design issue: functional side effects. This issue is discussed in Section 9.10.

9.5.8 Examples of Parameter Passing

Consider the following C function:

```c
void swap1(int a, int b) {
    int temp = a;
    a = b;
    b = temp;
}
```
Suppose this function is called with

`swap1(c, d);`

Recall that C uses pass-by-value. The actions of `swap1` can be described by the following pseudocode:

```c
a = c        - Move first parameter value in
b = d        - Move second parameter value in
temp = a
a = b
b = temp
```

Although `a` ends up with `d`’s value and `b` ends up with `c`’s value, the values of `c` and `d` are unchanged because nothing is transmitted back to the caller.

We can modify the C swap function to deal with pointer parameters to achieve the effect of pass-by-reference:

```c
void swap2(int *a, int *b) {
    int temp = *a;
    *a = *b;
    *b = temp;
}
```

`swap2` can be called with

`swap2(&c, &d);`

The actions of `swap2` can be described with the following:

```c
a = &c       - Move first parameter address in
b = &d       - Move second parameter address in
temp = *a
*a = *b
*b = temp
```

In this case, the swap operation is successful: The values of `c` and `d` are in fact interchanged. `swap2` can be written in C++ using reference parameters as follows:

```cpp
void swap2(int &a, int &b) {
    int temp = a;
    a = b;
    b = temp;
}
```

This simple swap operation is not possible in Java, because it has neither pointers nor C++’s kind of references. In Java, a reference variable can point to only an object, not a scalar value.
The semantics of pass-by-value-result is identical to those of pass-by-reference, except when aliasing is involved. Ada uses pass-by-value-result for inout-mode scalar parameters. To explore pass-by-value-result, consider the following function, `swap3`, which we assume uses pass-by-value-result parameters. It is written in a syntax similar to that of Ada.

```plaintext
procedure swap3(a : in out Integer, b : in out Integer) is
  temp : Integer;
  begin
    temp := a;
    a := b;
    b := temp;
  end swap3;
```

Suppose `swap3` is called with

```
swap3(c, d);
```

The actions of `swap3` with this call are

- `addr_c = &c` - Move first parameter address in
- `addr_d = &d` - Move second parameter address in
- `a = *addr_c` - Move first parameter value in
- `b = *addr_d` - Move second parameter value in
- `temp = a`
- `a = b`
- `b = temp`
- `*addr_c = a` - Move first parameter value out
- `*addr_d = b` - Move second parameter value out

So once again, this swap subprogram operates correctly. Next, consider the call

```
swap3(i, list[i]);
```

In this case, the actions are

- `addr_i = &i` - Move first parameter address in
- `addr_listi = &list[i]` - Move second parameter address in
- `a = *addr_i` - Move first parameter value in
- `b = *addr_listi` - Move second parameter value in
- `temp = a`
- `a = b`
- `b = temp`
- `*addr_i = a` - Move first parameter value out
- `*addr_listi = b` - Move second parameter value out

Again, the subprogram operates correctly, in this case because the addresses to which to return the values of the parameters are computed at the time of the call rather than at the time of the return. If the addresses of the actual parameters were computed at the time of the return, the results would be wrong.
Finally, we must explore what happens when aliasing is involved with pass-by-value-result and pass-by-reference. Consider the following skeletal program written in C-like syntax:

```c
int i = 3; /* i is a global variable */
void fun(int a, int b) {
    i = b;
}
void main() {
    int list[10];
    list[i] = 5;
    fun(i, list[i]);
}
```

In `fun`, if pass-by-reference is used, `i` and `a` are aliases. If pass-by-value-result is used, `i` and `a` are not aliases. The actions of `fun`, assuming pass-by-value-result, are as follows:

- `addr_i = &i` - Move first parameter address in
- `addr_listi = &list[i]` - Move second parameter address in
- `a = *addr_i` - Move first parameter value in
- `b = *addr_listi` - Move second parameter value in
- `i = b` - Sets `i` to 5
- `*addr_i = a` - Move first parameter value out
- `*addr_listi = b` - Move second parameter value out

In this case, the assignment to the global `i` in `fun` changes its value from 3 to 5, but the copy back of the first formal parameter (the second to last line in the example) sets it back to 3. The important observation here is that if pass-by-reference is used, the result is that the copy back is not part of the semantics, and `i` remains 5. Also note that because the address of the second parameter is computed at the beginning of `fun`, any change to the global `i` has no effect on the address used at the end to return the value of `list[i]`.

### 9.6 Parameters That Are Subprograms

In programming, a number of situations occur that are most conveniently handled if subprogram names can be sent as parameters to other subprograms. One common example of these occurs when a subprogram must sample some mathematical function. For example, a subprogram that does numerical integration estimates the area under the graph of a function by sampling the function at a number of different points. When such a subprogram is written, it should be usable for any given function; it should not need to be rewritten for every function that must be integrated. It is therefore natural that the name of a program function that evaluates the mathematical function to be integrated be sent to the integrating subprogram as a parameter.
Although the idea is natural and seemingly simple, the details of how it works can be confusing. If only the transmission of the subprogram code was necessary, it could be done by passing a single pointer. However, two complications arise.

First, there is the matter of type checking the parameters of the activations of the subprogram that was passed as a parameter. In C and C++, functions cannot be passed as parameters, but pointers to functions can. The type of a pointer to a function includes the function’s protocol. Because the protocol includes all parameter types, such parameters can be completely type checked.

The second complication with parameters that are subprograms appears only with languages that allow nested subprograms. The issue is what referencing environment for executing the passed subprogram should be used. There are three choices:

- The environment of the call statement that enacts the passed subprogram (shallow binding)
- The environment of the definition of the passed subprogram (deep binding)
- The environment of the call statement that passed the subprogram as an actual parameter (ad hoc binding)

The following example program, written with the syntax of JavaScript, illustrates these choices:

```javascript
function sub1() {
  var x;
  function sub2() {
    alert(x);  // Creates a dialog box with the value of x
  }
  function sub3() {
    var x;
    x = 3;
    sub4(sub2);
  }
  function sub4(subx) {
    var x;
    x = 4;
    subx();
  }
  x = 1;
  sub3();
}
```

Consider the execution of `sub2` when it is called in `sub4`. For shallow binding, the referencing environment of that execution is that of `sub4`, so the reference to `x` in `sub2` is bound to the local `x` in `sub4`, and the output of the program is 4. For deep binding, the referencing environment of `sub2`’s execution is that of `sub1`, so the reference to `x` in `sub2` is bound to the local `x` in `sub1`, and the output is 1. For ad hoc binding, the binding is to the local `x` in `sub3`, and the output is 3.
9.7 Calling Subprograms Indirectly

There are situations in which subprograms must be called indirectly. These most often occur when the specific subprogram to be called is not known until run time. The call to the subprogram is made through a pointer or reference to the subprogram, which has been set during execution before the call is made. The two most common applications of indirect subprogram calls are for event handling in graphical user interfaces, which are now part of nearly all Web applications, as well as many non-Web applications, and for callbacks, in which a subprogram is called and instructed to notify the caller when the called subprogram has completed its work. As always, our interest is not in these specific kinds of programming, but rather in programming language support for them.

The concept of calling subprograms indirectly is not a recently developed concept. C and C++ allow a program to define a pointer to a function, through which the function can be called. In C++, pointers to functions are typed according to the return type and parameter types of the function, so that such a pointer can point only at functions with one particular protocol. For example, the following declaration defines a pointer \( pfun \) that can point to any function that takes a \texttt{float} and an \texttt{int} as parameters and returns a \texttt{float}:

\[
\text{float } (*pfun)(\text{float, int});
\]

Any function with the same protocol as this pointer can be used as the initial value of this pointer or be assigned to the pointer in a program. In C and C++, a function name without following parentheses, like an array name without
following brackets, is the address of the function (or array). So, both of the following are legal ways of giving an initial value or assigning a value to a pointer to a function:

```c
int myfun2 (int, int);  // A function declaration
int (*pfun2)(int, int) = myfun2;  // Create a pointer and initialize
                           // it to point to myfun2
pfun2 = myfun2;  // Assigning a function's address to a pointer
```

The function `myfun2` can now be called with either of the following statements:

```c
(*pfun2)(first, second); pfun2(first, second);
```

The first of these explicitly dereferences the pointer `pfun2`, which is legal, but unnecessary.

The function pointers of C and C++ can be sent as parameters and returned from functions, although functions cannot be used directly in either of those roles.

In C#, the power and flexibility of method pointers is increased by making them objects. These are called delegates, because instead of calling a method, a program delegates that action to a delegate.

To use a delegate, first the delegate class must be defined with a specific method protocol. An instantiation of a delegate holds the name of a method with the delegate's protocol that it is able to call. The syntax of a declaration of a delegate is the same as that of a method declaration, except that the reserved word `delegate` is inserted just before the return type. For example, we could have the following:

```c
public delegate int Change(int x);
```

This delegate can be instantiated with any method that takes an `int` as a parameter and returns an `int`. For example, consider the following method declaration:

```c
static int fun1(int x);
```

The delegate `Change` can be instantiated by sending the name of this method to the delegate's constructor, as in the following:

```c
Change chgfun1 = new Change(fun1);
```

This can be shortened to the following:

```c
Change chgfun1 = fun1;
```
Following is an example call to `fun1` through the delegate `chgfun1`:

```c
chgfun1(12);
```

Objects of a delegate class can store more than one method. A second method can be added using the operator `+=`, as in the following:

```c
Change chgfun1 += fun2;
```

This places `fun2` in the `chgfun1` delegate, even if `chgfun1` previously had the value `null`. All of the methods stored in a delegate instance are called in the order in which they were placed in the instance. This is called a **multicast delegate**. Regardless of what is returned by the methods, only the value or object returned by the last one called is returned. Of course, this means that in most cases, `void` is returned by the methods called through a multicast delegate.

In our example, a static method is placed in the delegate `Change`. Instance methods can also be called through a delegate, in which case the delegate must store a reference to the method. Delegates can also be generic.

Delegates are used for event handling by .NET applications. They are also used to implement closures (see Section 9.12).

As is the case with C and C++, the name of a function in Python without the following parentheses is a pointer to that function. Ada 95 has pointers to subprograms, but Java does not. In Python and Ruby, as well as most functional languages, subprograms are treated like data, so they can be assigned to variables. Therefore, in these languages, there is little need for pointers to subprograms.

### 9.8 Design Issues for Functions

The following design issues are specific to functions:

- Are side effects allowed?
- What types of values can be returned?
- How many values can be returned?

#### 9.8.1 Functional Side Effects

Because of the problems of side effects of functions that are called in expressions, as described in Chapter 5, parameters to functions should always be in-mode. In fact, some languages require this; for example, Ada functions can have only in-mode formal parameters. This requirement effectively prevents a function from causing side effects through its parameters or through aliasing of parameters and globals. In most other imperative languages, however, functions
can have either pass-by-value or pass-by-reference parameters, thus allowing functions that cause side effects and aliasing.

Pure functional languages, such as Haskell, do not have variables, so their functions cannot have side effects.

### 9.8.2 Types of Returned Values

Most imperative programming languages restrict the types that can be returned by their functions. C allows any type to be returned by its functions except arrays and functions. Both of these can be handled by pointer type return values. C++ is like C but also allows user-defined types, or classes, to be returned from its functions. Ada, Python, Ruby, and Lua are the only languages among current imperative languages whose functions (and/or methods) can return values of any type. In the case of Ada, however, because functions are not types in Ada, they cannot be returned from functions. Of course, pointers to functions can be returned by functions.

In some programming languages, subprograms are first-class objects, which means that they can be passed as parameters, returned from functions, and assigned to variables. Methods are first-class objects in some imperative languages, for example, Python, Ruby, and Lua. The same is true for the functions in most functional languages.

Neither Java nor C# can have functions, although their methods are similar to functions. In both, any type or class can be returned by methods. Because methods are not types, they cannot be returned.

### 9.8.3 Number of Returned Values

In most languages, only a single value can be returned from a function. However, that is not always the case. Ruby allows the return of more than one value from a method. If a `return` statement in a Ruby method is not followed by an expression, `nil` is returned. If followed by one expression, the value of the expression is returned. If followed by more than one expression, an array of the values of all of the expressions is returned.

Lua also allows functions to return multiple values. Such values follow the `return` statement as a comma-separated list, as in the following:

```lua
return 3, sum, index
```

The form of the statement that calls the function determines the number of values that are received by the caller. If the function is called as a procedure, that is, as a statement, all return values are ignored. If the function returned three values and all are to be kept by the caller, the function would be called as in the following example:

```plaintext
a, b, c = fun()
```

In F#, multiple values can be returned by placing them in a tuple and having the tuple be the last expression in the function.
9.9 Overloaded Subprograms

An overloaded operator is one that has multiple meanings. The meaning of a particular instance of an overloaded operator is determined by the types of its operands. For example, if the * operator has two floating-point operands in a Java program, it specifies floating-point multiplication. But if the same operator has two integer operands, it specifies integer multiplication.

An overloaded subprogram is a subprogram that has the same name as another subprogram in the same referencing environment. Every version of an overloaded subprogram must have a unique protocol; that is, it must be different from the others in the number, order, or types of its parameters, and possibly in its return type. The meaning of a call to an overloaded subprogram is determined by the actual parameter list and/or possibly the type of the returned value. Although it is not necessary, overloaded subprograms usually implement the same process.

C++, Java, and C# include predefined overloaded subprograms. For example, many classes in C++, Java, and C# have overloaded constructors. Because each version of an overloaded subprogram has a unique parameter profile, the compiler can disambiguate occurrences of calls to them by the different type parameters. Unfortunately, it is not that simple. Parameter coercions, when allowed, complicate the disambiguation process enormously. Simply stated, the issue is that if no method’s parameter profile matches the number and types of the actual parameters in a method call, but two or more methods have parameter profiles that can be matched through coercions, which method should be called? For a language designer to answer this question, he or she must decide how to rank all of the different coercions, so that the compiler can choose the method that “best” matches the call. This can be a complicated task. To understand the level of complexity of this process, we suggest the reader refer to the rules for disambiguation of method calls used in C++ (Stroustrup, 1997).

Because C++, Java, and C# allow mixed-mode expressions, the return type is irrelevant to disambiguation of overloaded functions (or methods). The context of the call does not allow the determination of the return type. For example, if a C++ program has two functions named **fun** and both take an int parameter but one returns an int and one returns a float, the program would not compile, because the compiler could not determine which version of **fun** should be used.

Users are also allowed to write multiple versions of subprograms with the same name in Java, C++, C#, and F#. Once again, in C++, Java, and C# the most common user-defined overloaded methods are constructors.

Overloaded subprograms that have default parameters can lead to ambiguous subprogram calls. For example, consider the following C++ code:

```cpp
void fun(float b = 0.0);
void fun();
....
fun();
```

The call is ambiguous and will cause a compilation error.
Software reuse can be an important contributor to software productivity. One way to increase the reusability of software is to lessen the need to create different subprograms that implement the same algorithm on different types of data. For example, a programmer should not need to write four different sort subprograms to sort four arrays that differ only in element type.

A polymorphic subprogram takes parameters of different types on different activations. Overloaded subprograms provide a particular kind of polymorphism called ad hoc polymorphism. Overloaded subprograms need not behave similarly.

Languages that support object-oriented programming usually support subtype polymorphism. Subtype polymorphism means that a variable of type T can access any object of type T or any type derived from T.

A more general kind of polymorphism is provided by the methods of Python and Ruby. Recall that variables in these languages do not have types, so formal parameters do not have types. Therefore, a method will work for any type of actual parameter, as long as the operators used on the formal parameters in the method are defined.

Parametric polymorphism is provided by a subprogram that takes generic parameters that are used in type expressions that describe the types of the parameters of the subprogram. Different instantiations of such subprograms can be given different generic parameters, producing subprograms that take different types of parameters. Parametric definitions of subprograms all behave the same. Parametrically polymorphic subprograms are often called generic subprograms. C++, Java 5.0+, C# 2005+, and F# provide a kind of compile-time parametric polymorphism.

### 9.10.1 Generic Functions in C++

Generic functions in C++ have the descriptive name of template functions. The definition of a template function has the general form

```
template <template parameters>
— a function definition that may include the template parameters
```

A template parameter (there must be at least one) has one of the forms

```
class identifier
typename identifier
```

The class form is used for type names. The typename form is used for passing a value to the template function. For example, it is sometimes convenient to pass an integer value for the size of an array in the template function.
A template can take another template, in practice often a template class that defines a user-defined generic type, as a parameter, but we do not consider that option here.\(^8\)

As an example of a template function, consider the following:

```cpp
template <class Type>
Type max(Type first, Type second) {
    return first > second ? first : second;
}
```

where `Type` is the parameter that specifies the type of data on which the function will operate. This template function can be instantiated for any type for which the operator `>` is defined. For example, if it were instantiated with `int` as the parameter, it would be

```cpp
int max(int first, int second) {
    return first > second ? first : second;
}
```

Although this process could be defined as a macro, a macro would have the disadvantage of not operating correctly if the parameters were expressions with side effects. For example, suppose the macro were defined as

```cpp
#define max(a, b) ((a) > (b)) ? (a) : (b)
```

This definition is generic in the sense that it works for any numeric type. However, it does not always work correctly if called with a parameter that has a side effect, such as

```cpp
max(x++, y)
```

which produces

```cpp
((x++) > (y) ? (x++) : (y))
```

Whenever the value of `x` is greater than that of `y`, `x` will be incremented twice.

C++ template functions are instantiated implicitly either when the function is named in a call or when its address is taken with the `&` operator. For example, the example template function `max` would be instantiated twice by the following code segment—one for `int` type parameters and once for `char` type parameters:

```cpp
int a, b, c;
char d, e, f;
```

---

8. Template classes are discussed in Chapter 11.
The following is a C++ generic sort subprogram:

```cpp
template <class Type>
void generic_sort(Type list[], int len) {
    int top, bottom;
    Type temp;
    for (top = 0; top < len - 2; top++)
        for (bottom = top + 1; bottom < len - 1; bottom++)
            if (list[top] > list[bottom]) {
                temp = list[top];
                list[top] = list[bottom];
                list[bottom] = temp;
            }  //** end of if (list[top] ... 
}  //** end of generic_sort
```

The following is an example instantiation of this template function:

```cpp
float flt_list[100];
... 
generic_sort(flt_list, 100);
```

The templated functions of C++ are a kind of poor cousin to a subprogram in which the types of the formal parameters are dynamically bound to the types of the actual parameters in a call. In this case, only a single copy of the code is needed, whereas with the C++ approach, a copy must be created at compile time for each different type that is required and the binding of subprogram calls to subprograms is static.

### 9.10.2 Generic Methods in Java 5.0

Support for generic types and methods was added to Java in Java 5.0. The name of a generic class in Java 5.0 is specified by a name followed by one or more type variables delimited by pointed brackets. For example,

```java
generic_class<T>
```

where `T` is the type variable. Generic types are discussed in more detail in Chapter 11.

Java’s generic methods differ from the generic subprograms of C++ in several important ways. First, generic parameters must be classes—they cannot be primitive types. This requirement disallows a generic method that mimics our example in C++, in which the component types of arrays are generic and can be primitives. In Java, the components of arrays (as opposed to containers) cannot be generic. Second, although Java generic methods can be instantiated...
any number of times, only one copy of the code is built. The internal version of a generic method, which is called a raw method, operates on Object class objects. At the point where the generic value of a generic method is returned, the compiler inserts a cast to the proper type. Third, in Java, restrictions can be specified on the range of classes that can be passed to the generic method as generic parameters. Such restrictions are called bounds.

As an example of a generic Java 5.0 method, consider the following skeletal method definition:

```java
public static <T> T doIt(T[] list) {
    ...
}
```

This defines a method named doIt that takes an array of elements of a generic type. The name of the generic type is T and it must be an array. Following is an example call to doIt:

```java
doit<String>(myList);
```

Now, consider the following version of doIt, which has a bound on its generic parameter:

```java
public static <T extends Comparable> T doIt(T[] list) {
    ...
}
```

This defines a method that takes a generic array parameter whose elements are of a class that implements the Comparable interface. That is the restriction, or bound, on the generic parameter. The reserved word extends seems to imply that the generic class subclasses the following class. In this context, however, extends has a different meaning. The expression <T extends BoundingType> specifies that T should be a “subtype” of the bounding type. So, in this context, extends means the generic class (or interface) either extends the bounding class (the bound if it is a class) or implements the bounding interface (if the bound is an interface). The bound ensures that the elements of any instantiation of the generic can be compared with the Comparable method, compareTo.

If a generic method has two or more restrictions on its generic type, they are added to the extends clause, separated by ampersands (&). Also, generic methods can have more than one generic parameter.

Java 5.0 supports wildcard types. For example, Collection<?> is a wildcard type for collection classes. This type can be used for any collection type of any class components. For example, consider the following generic method:

```java
void printCollection(Collection<?> c) {
    for (Object e: c) {
        System.out.println(e);
    }
}
```
This method prints the elements of any `Collection` class, regardless of the class of its components. Some care must be taken with objects of the wildcard type. For example, because the components of a particular object of this type have a type, other type objects cannot be added to the collection. For example, consider

```java
Collection<?> c = new ArrayList<String>();
```

It would be illegal to use the `add` method to put something into this collection unless its type were `String`.

Wildcard types can be restricted, as is the case with nonwildcard types. Such types are called *bounded wildcard types*. For example, consider the following method header:

```java
public void drawAll(ArrayList<? extends Shape> things)
```

The generic type here is a wildcard type that is a subclass of the `Shape` class. This method could be written to draw any object whose type is a subclass of `Shape`.

### 9.10.3 Generic Methods in C# 2005

The generic methods of C# 2005 are similar in capability to those of Java 5.0, except there is no support for wildcard types. One unique feature of C# 2005 generic methods is that the actual type parameters in a call can be omitted if the compiler can infer the unspecified type. For example, consider the following skeletal class definition:

```csharp
class MyClass {
    public static T DoIt<T>(T p1) {
        ...
    }
}
```

The method `DoIt` can be called without specifying the generic parameter if the compiler can infer the generic type from the actual parameter in the call. For example, both of the following calls are legal:

```csharp
int myInt = MyClass.DoIt(17); // Calls DoIt<int>
string myStr = MyClass.DoIt('apples'); // Calls DoIt<string>
```

### 9.10.4 Generic Functions in F#

The type inferencing system of F# is not always able to determine the type of parameters or the return type of a function. When this is the case, for some functions F# infers a generic type for the parameters and the return value.
This is called **automatic generalization**. For example, consider the following function definition:

```fsharp
let getLast (a, b, c) = c;;
```

Because no type information was included, the types of the parameters and the return value are all inferred to be generic. Because this function does not include any computations, this is a simple generic function.

Functions can be defined to have generic parameters, as in the following example:

```fsharp
let printPair (x: 'a) (y: 'a) =
    printfn "%A %A" x y;;
```

The `%A` format specification is for any type. The apostrophe in front of the type named `a` specifies it to be a generic type. This function definition works (with generic parameters) because no type-constrained operation is included. Arithmetic operators are examples of type-constrained operations. For example, consider the following function definition:

```fsharp
let adder x y = x + y;;
```

Type inferencing sets the type of `x` and `y` and the return value to `int`. Because there is no type coercion in F#, the following call is illegal:

```fsharp
adder 2.5 3.6;;
```

Even if the type of the parameters were set to be generic, the `+` operator would cause the types of `x` and `y` to be `int`.

The generic type could also be specified explicitly in angle brackets, as in the following:

```fsharp
let printPair2<'T> x y =
    printfn "%A %A" x y;;
```

This function must be called with a type, as in the following:

```fsharp
printPair2<float> 3.5 2.4;;
```

Because of type inferencing and the lack of type coercions, F# generic functions are far less useful, especially for numeric computations, than those of C++, Java 5.0+, and C# 2005+.

---

9. There is nothing special about `a`—it could be any legal identifier. By convention, lowercase letters at the beginning of the alphabet are used.

10. Convention explicitly states that generic types are named with uppercase letters starting at `T`. 
9.11 User-Defined Overloaded Operators

Operators can be overloaded by the user in Ada, C++, Python, and Ruby. Suppose that a Python class is developed to support complex numbers and arithmetic operations on them. A complex number can be represented with two floating-point values. The Complex class would have members for these two named real and imag. In Python, binary arithmetic operations are implemented as method calls sent to the first operand, sending the second operand as a parameter. For addition, the method is named __add__. For example, the expression $x + y$ is implemented as $x.__add__(y)$. To overload $+$ for the addition of objects of the new Complex class, we only need to provide Complex with a method named __add__ that performs the operation. Following is such a method:

```python
def __add__(self, second):
    return Complex(self.real + second.real, self.imag + second.imag)
```

In most languages that support object-oriented programming, a reference to the current object is implicitly sent with each method call. In Python, this reference must be sent explicitly; that is the reason why self is the first parameter to our method, __add__.

The example add method could be written for a complex class in C++ as follows:

```c++
Complex operator +(Complex &second) {
    return Complex(real + second.real, imag + second.imag);
}
```

9.12 Closures

Defining a closure is a simple matter; a closure is a subprogram and the referencing environment where it was defined. The referencing environment is needed if the subprogram can be called from any arbitrary place in the program. Explaining a closure is not so simple.

If a static-scoped programming language does not allow nested subprograms, closures are not useful, so such languages do not support them. All of the variables in the referencing environment of a subprogram in such a language (its local variables and the global variables) are accessible, regardless of the place in the program where the subprogram is called.

When subprograms can be nested, in addition to locals and globals, the referencing environment of a subprogram can include variables defined in all enclosing subprograms. However, this is not an issue if the subprogram can be

---

11. Both C++ and Python have predefined classes for complex numbers, so our example methods are unnecessary, except as illustrations.
called only in places where all of the enclosing scopes are active and visible. It becomes an issue if a subprogram can be called elsewhere. This can happen if the subprogram can be passed as a parameter or assigned to a variable, thereby allowing it to be called from virtually anywhere in the program. There is an associated problem: The subprogram could be called after one or more of its nesting subprograms has terminated, which normally means that the variables defined in such nesting subprograms have been deallocated—they no longer exist. For the subprogram to be callable from anywhere in the program, its referencing environment must be available wherever it might be called. Therefore, the variables defined in nesting subprograms may need lifetimes that are of the entire program, rather than just the time during which the subprogram in which they were defined is active. A variable whose lifetime is that of the whole program is said to have **unlimited extent**. This usually means they must be heap dynamic, rather than stack dynamic.

Nearly all functional programming languages, most scripting languages, and at least one primarily imperative language, C#, support closures. These languages are static-scoped, allow nested subprograms, and allow subprograms to be passed as parameters. Following is an example of a closure written in JavaScript:

```javascript
function makeAdder(x) {
    return function(y) { return x + y; }
}

var add10 = makeAdder(10);
var add5 = makeAdder(5);
document.write("Add 10 to 20: " + add10(20) + "<br />");
document.write("Add 5 to 20: " + add5(20) + "<br />");
```

The output of this code, assuming it was embedded in an HTML document and displayed with a browser, is as follows:

```
Add 10 to 20: 30
Add 5 to 20: 25
```

In this example, the closure is the anonymous function defined inside the `makeAdder` function, which `makeAdder` returns. The variable `x` referenced in the closure function is bound to the parameter that was sent to `makeAdder`. The `makeAdder` function is called twice, once with a parameter of 10 and once with 5. Each of these calls returns a different version of the closure because they are bound to different values of `x`. The first call to `makeAdder` creates a function that adds 10 to its parameter; the second creates a function that adds 5 to its parameter. The two versions of the function are bound to

---

12. In C#, the only methods that can be nested are anonymous delegates and lambda expressions.
different activations of makeAdder. Obviously, the lifetime of the version of x created when makeAdder is called must extend over the lifetime of the program.

This same closure function can be written in C# using a nested anonymous delegate. The type of the nesting method is specified to be a function that takes an int as a parameter and returns an anonymous delegate. The return type is specified with the special notation for such delegates, Func<int, int>. The first type in the angle brackets is the parameter type. Such a delegate can encapsulate methods that have only one parameter. The second type is the return type of the method encapsulated by the delegate.

```csharp
static Func<int, int> makeAdder(int x) {
    return delegate(int y) { return x + y; };
}
```

```csharp
Func<int, int> Add10 = makeAdder(10);
Func<int, int> Add5 = makeAdder(5);
Console.WriteLine("Add 10 to 20: {0}", Add10(20));
Console.WriteLine("Add 5 to 20: {0}", Add5(20));
```

The output of this code is exactly the same as for the previous JavaScript closure example.

The anonymous delegate could have been written as a lambda expression. The following is a replacement for the body of the makeAdder method, using a lambda expression instead of the delegate:

```csharp
return y => x + y
```

Ruby’s blocks are implemented so that they can reference variables visible in the position in which they were defined, even if they are called at a place in which those variables would have disappeared. This makes such blocks closures.

### 9.13 Coroutines

A coroutine is a special kind of subprogram. Rather than the master-slave relationship between a caller and a called subprogram that exists with conventional subprograms, caller and called coroutines are more equitable. In fact, the coroutine control mechanism is often called the symmetric unit control model.

Coroutines can have multiple entry points, which are controlled by the coroutines themselves. They also have the means to maintain their status between activations. This means that coroutines must be history sensitive and thus have static local variables. Secondary executions of a coroutine often begin at points other than its beginning. Because of this, the invocation of a coroutine is called a resume rather than a call.
For example, consider the following skeletal coroutine:

```plaintext
sub co1() {
  . . .
  resume co2();
  . . .
  resume co3();
  . . .
}
```

The first time `co1` is resumed, its execution begins at the first statement and executes down to and including the resume of `co2`, which transfers control to `co2`. The next time `co1` is resumed, its execution begins at the first statement after its call to `co2`. When `co1` is resumed the third time, its execution begins at the first statement after the resume of `co3`.

One of the usual characteristics of subprograms is maintained in coroutines: Only one coroutine is actually in execution at a given time.

As seen in the example above, rather than executing to its end, a coroutine often partially executes and then transfers control to some other coroutine, and when restarted, a coroutine resumes execution just after the statement it used to transfer control elsewhere. This sort of interleaved execution sequence is related to the way multiprogramming operating systems work. Although there may be only one processor, all of the executing programs in such a system appear to run concurrently while sharing the processor. In the case of coroutines, this is sometimes called quasi-concurrency.

Typically, coroutines are created in an application by a program unit called the master unit, which is not a coroutine. When created, coroutines execute their initialization code and then return control to that master unit. When the entire family of coroutines is constructed, the master program resumes one of the coroutines, and the members of the family of coroutines then resume each other in some order until their work is completed, if in fact it can be completed. If the execution of a coroutine reaches the end of its code section, control is transferred to the master unit that created it. This is the mechanism for ending execution of the collection of coroutines, when that is desirable. In some programs, the coroutines run whenever the computer is running.

One example of a problem that can be solved with this sort of collection of coroutines is a card game simulation. Suppose the game has four players who all use the same strategy. Such a game can be simulated by having a master program unit create a family of four coroutines, each with a collection, or hand, of cards. The master program could then start the simulation by resuming one of the player coroutines, which, after it had played its turn, could resume the next player coroutine, and so forth until the game ended.

Suppose program units `A` and `B` are coroutines. Figure 9.3 shows two ways an execution sequence involving `A` and `B` might proceed.

In Figure 9.3a, the execution of coroutine `A` is started by the master unit. After some execution, `A` starts `B`. When coroutine `B` in Figure 9.3a first
Figure 9.3
Two possible execution control sequences for two coroutines without loops

Figure 9.4
Coroutine execution sequence with loops

causes control to return to coroutine A, the semantics is that A continues from where it ended its last execution. In particular, its local variables have the values left them by the previous activation. Figure 9.3b shows an alternative execution sequence of coroutines A and B. In this case, B is started by the master unit.
Rather than have the patterns shown in Figure 9.3, a coroutine often has a loop containing a resume. Figure 9.4 shows the execution sequence of this scenario. In this case, \( A \) is started by the master unit. Inside its main loop, \( A \) resumes \( B \), which in turn resumes \( A \) in its main loop.

Among contemporary languages, only Lua fully supports coroutines.\(^{13}\)

Summary

Process abstractions are represented in programming languages by subprograms. A subprogram definition describes the actions represented by the subprogram. A subprogram call enacts those actions. A subprogram header identifies a subprogram definition and provides its interface, which is called its protocol.

Formal parameters are the names that subprograms use to refer to the actual parameters given in subprogram calls. In Python and Ruby, array and hash formal parameters are used to support variable numbers of parameters. Lua and JavaScript also support variable numbers of parameters. Actual parameters can be associated with formal parameters by position or by keyword. Parameters can have default values.

Subprograms can be either functions, which model mathematical functions and are used to define new operations, or procedures, which define new statements.

Local variables in subprograms can be stack dynamic, providing support for recursion, or static, providing efficiency and history-sensitive local variables.

JavaScript, Python, Ruby, and Lua allow subprogram definitions to be nested.

There are three fundamental semantics models of parameter passing—in mode, out mode, and inout mode—and a number of approaches to implement them. These are pass-by-value, pass-by-result, pass-by-value-result, pass-by-reference, and pass-by-name. In most languages, parameters are passed in the run-time stack.

Aliasing can occur when pass-by-reference parameters are used, both among two or more parameters and between a parameter and an accessible nonlocal variable.

Parameters that are multidimensioned arrays pose some issues for the language designer, because the called subprogram needs to know how to compute the storage mapping function for them. This requires more than just the name of the array.

Parameters that are subprogram names provide a necessary service but can be difficult to understand. The opacity lies in the referencing environment that is available when a subprogram that has been passed as a parameter is executed.

C and C++ support pointers to functions. C# has delegates, which are objects that can store references to methods. Delegates can support multicast calls by storing more than one method reference.

---

\(^{13}\) However, the generators of Python are a form of coroutines.
Ada, C++, C#, Ruby, and Python allow both subprogram and operator overloading. Subprograms can be overloaded as long as the various versions can be disambiguated by the types of their parameters or returned values. Function definitions can be used to build additional meanings for operators.

Subprograms in C++, Java 5.0, and C# 2005 can be generic, using parametric polymorphism, so the desired types of their data objects can be passed to the compiler, which then can construct units for the requested types.

The designer of a function facility in a language must decide what restrictions will be placed on the returned values, as well as the number of return values.

A closure is a subprogram and its referencing environment. Closures are useful in languages that allow nested subprograms, are static-scoped, and allow subprograms to be returned from functions and assigned to variables.

A coroutine is a special subprogram that has multiple entries. It can be used to provide interleaved execution of subprograms.

**REVIEW QUESTIONS**

1. What are the three general characteristics of subprograms?
2. What is a subprogram call?
3. What is a subprogram definition?
4. What characteristic of Python subprograms sets them apart from those of other languages?
5. How do Ruby methods differ from the subprograms of other programming languages?
6. What is the feature of Lua functions?
7. What are function declarations called in C and C++? Where are the declarations often placed?
8. Name one pure functional programming language that does not have mutable data.
9. What are positional parameters?
10. Give an example of a language that allows positional parameters in addition to keyword parameters.
11. What is the use of a default value in a formal parameter?
12. What is the rule of using a default parameter in C++?
13. What is the rule for accepting variable parameters in C# methods?
14. What language allows array formal parameters?
15. What is an ellipsis?
16. What are the modes, the conceptual models of transfer, the advantages, and the disadvantages of pass-by-value, pass-by-result, pass-by-value-result, and pass-by-reference parameter-passing methods?
17. Describe the ways that aliases can occur with pass-by-reference parameters.
18. What is the difference between the way original C and C89 deal with an actual parameter whose type is not identical to that of the corresponding formal parameter?

19. Name some languages that support procedures.

20. Describe the problem of passing multidimensioned arrays as parameters.

21. What is the name of the parameter-passing method used in Ruby?

22. What are the two issues that arise when subprogram names are parameters?

23. Define shallow and deep binding for referencing environments of subprograms that have been passed as parameters.

24. What is a generic subprogram?

25. What is ad hoc binding?

26. What causes a C++ template function to be instantiated?

27. In what fundamental ways do the generic methods of Java 5.0 differ from those of C# 2005?

28. If a Java 5.0 method returns a generic type, what type of object is actually returned?

29. If a Java 5.0 generic method is called with three different generic parameters, how many versions of the method will be generated by the compiler?

30. When does a variable have unlimited extent?

31. What is subtype polymorphism?

32. What is a multicast delegate?

33. What is the main drawback of generic functions in F#?

34. What is a coroutine?

35. What are the language characteristics that make closures useful?

36. What languages allow the user to overload operators?

37. What is a symmetric unit control model?

---

**Problem Set**

1. What are arguments for and against a user program building additional definitions for existing operators, as can be done in Python and C++? Do you think such user-defined operator overloading is good or bad? Support your answer.
2. In most Fortran IV implementations, all parameters were passed by reference, using access path transmission only. State both the advantages and disadvantages of this design choice.

3. Argue in support of the Ada 83 designers’ decision to allow the implementor to choose between implementing \texttt{inout}-mode parameters by copy or by reference.

4. Suppose you want to write a method that prints a heading on a new output page, along with a page number that is 1 in the first activation and that increases by 1 with each subsequent activation. Can this be done without parameters and without reference to nonlocal variables in Java? Can it be done in C#?

5. Consider the following program written in C syntax:

```c
void swap(int a, int b) {
    int temp;
    temp = a;
    a = b;
    b = temp;
}
void main() {
    int value = 2, list[5] = {1, 3, 5, 7, 9};
    swap(value, list[0]);
    swap(list[0], list[1]);
    swap(value, list[value]);
}
```

For each of the following parameter-passing methods, what are all of the values of the variables \texttt{value} and \texttt{list} after each of the three calls to \texttt{swap}?

a. Passed by value  
b. Passed by reference  
c. Passed by value-result

6. Present one argument against providing both static and dynamic local variables in subprograms.

7. Consider the following program written in C syntax:

```c
void fun (int first, int second) {
    first += first;
    second += second;
}
void main() {
    int list[2] = {1, 3};
    fun(list[0], list[1]);
}
```

For each of the following parameter-passing methods, what are the values of the \texttt{list} array after execution?
a. Passed by value  
b. Passed by reference  
c. Passed by value-result  

8. Argue against the C design of providing only function subprograms.  
9. From a textbook on Fortran, learn the syntax and semantics of statement functions. Justify their existence in Fortran.  
10. Study the methods of user-defined operator overloading in C++ and Ada, and write a report comparing the two using our criteria for evaluating languages.  
11. C# supports out-mode parameters, but neither Java nor C++ does. Explain the difference.  
12. Research Jensen’s Device, which was a use of pass-by-name parameters, and write a short description of what it is and how it can be used.  
13. Study the iterator mechanisms of Ruby and CLU and list their similarities and differences.  
14. Speculate on the issue of allowing nested subprograms in programming languages—why are they not allowed in many contemporary languages?  
15. What are at least two arguments against the use of pass-by-name parameters?  
16. Write a detailed comparison of the generic subprograms of Java 5.0 and C# 2005.  

**Programming Exercises**

1. Write a program in a language that you know to determine the ratio of the time required to pass a large array by reference and the time required to pass the same array by value. Make the array as large as possible on the machine and implementation you use. Pass the array as many times as necessary to get reasonably accurate timings of the passing operations.  
2. Write a C# or Ada program that determines when the address of an out-mode parameter is computed (at the time of the call or at the time the execution of the subprogram finishes).  
3. Write a Perl program that passes by reference a literal to a subprogram, which attempts to change the parameter. Given the overall design philosophy of Perl, explain the results.  
4. Repeat Programming Exercise 3 in C#.  
5. Write a program in some language that has both static and stack-dynamic local variables in subprograms. Create six large (at least 100 × 100) matrices in the subprogram—three static and three stack dynamic. Fill two of the static matrices and two of the stack-dynamic matrices with
random numbers in the range of 1 to 100. The code in the subprogram must perform a large number of matrix multiplication operations on the static matrices and time the process. Then it must repeat this with the stack-dynamic matrices. Compare and explain the results.

6. Write a C# program that includes two methods that are called a large number of times. Both methods are passed a large array, one by value and one by reference. Compare the times required to call these two methods and explain the difference. Be sure to call them a sufficient number of times to illustrate a difference in the required time.

7. Write a program, using the syntax of whatever language you like, that produces different behavior depending on whether pass-by-reference or pass-by-value-result is used in its parameter passing.

8. Write a generic C++ function that takes an array of generic elements and a scalar of the same type as the array elements. The type of the array elements and the scalar is the generic parameter. The function must search the given array for the given scalar and return the subscript of the scalar in the array. If the scalar is not in the array, the function must return -1. Test the function for int and float types.

9. Devise a subprogram and calling code in which pass-by-reference and pass-by-value-result of one or more parameters produces different results.
Implementing Subprograms

10.1 The General Semantics of Calls and Returns
10.2 Implementing “Simple” Subprograms
10.3 Implementing Subprograms with Stack-Dynamic Local Variables
10.4 Nested Subprograms
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Chapter 10   Implementing Subprograms

The purpose of this chapter is to explore the implementation of subprograms. The discussion will provide the reader with some knowledge of how subprogram linkage works, and also why ALGOL 60 was a challenge to the unsuspecting compiler writers of the early 1960s. We begin with the simplest situation, nonnestable subprograms with static local variables, advance to more complicated subprograms with stack-dynamic local variables, and conclude with nested subprograms with stack-dynamic local variables and static scoping. The increased difficulty of implementing subprograms in languages with nested subprograms is caused by the need to include mechanisms to access nonlocal variables.

The static chain method of accessing nonlocals in static-scoped languages is discussed in detail. Then, techniques for implementing blocks are described. Finally, several methods of implementing nonlocal variable access in a dynamic-scoped language are discussed.

10.1 The General Semantics of Calls and Returns

The subprogram call and return operations are together called subprogram linkage. The implementation of subprograms must be based on the semantics of the subprogram linkage of the language being implemented.

A subprogram call in a typical language has numerous actions associated with it. The call process must include the implementation of whatever parameter-passing method is used. If local variables are not static, the call process must allocate storage for the locals declared in the called subprogram and bind those variables to that storage. It must save the execution status of the calling program unit. The execution status is everything needed to resume execution of the calling program unit. This includes register values, CPU status bits, and the environment pointer (EP). The EP, which is further discussed in Section 10.3, is used to access parameters and local variables during the execution of a subprogram. The calling process also must arrange to transfer control to the code of the subprogram and ensure that control can return to the proper place when the subprogram execution is completed. Finally, if the language supports nested subprograms, the call process must create some mechanism to provide access to nonlocal variables that are visible to the called subprogram.

The required actions of a subprogram return are less complicated than those of a call. If the subprogram has parameters that are out mode or inout mode and are implemented by copy, the first action of the return process is to move the local values of the associated formal parameters to the actual parameters. Next, it must deallocate the storage used for local variables and restore the execution status of the calling program unit. Finally, control must be returned to the calling program unit.
10.2 Implementing “Simple” Subprograms

We begin with the task of implementing simple subprograms. By “simple” we mean that subprograms cannot be nested and all local variables are static. Early versions of Fortran were examples of languages that had this kind of subprograms.

The semantics of a call to a “simple” subprogram requires the following actions:

1. Save the execution status of the current program unit.
2. Compute and pass the parameters.
3. Pass the return address to the called.
4. Transfer control to the called.

The semantics of a return from a simple subprogram requires the following actions:

1. If there are pass-by-value-result or out-mode parameters, the current values of those parameters are moved to or made available to the corresponding actual parameters.
2. If the subprogram is a function, the functional value is moved to a place accessible to the caller.
3. The execution status of the caller is restored.
4. Control is transferred back to the caller.

The call and return actions require storage for the following:

- Status information about the caller
- Parameters
- Return address
- Return value for functions
- Temporaries used by the code of the subprograms

These, along with the local variables and the subprogram code, form the complete collection of information a subprogram needs to execute and then return control to the caller.

The question now is the distribution of the call and return actions to the caller and the called. For simple subprograms, the answer is obvious for most of the parts of the process. The last three actions of a call clearly must be done by the caller. Saving the execution status of the caller could be done by either. In the case of the return, the first, third, and fourth actions must be done by the called. Once again, the restoration of the execution status of the caller could be done by either the caller or the called. In general, the linkage actions of the called can occur at two different times, either at the beginning of its execution or at the end. These are sometimes called the prologue and epilogue of the subprogram linkage. In the case of a simple subprogram, all of the linkage actions of the callee occur at the end of its execution, so there is no need for a prologue.
A simple subprogram consists of two separate parts: the actual code of the subprogram, which is constant, and the local variables and data listed previously, which can change when the subprogram is executed. In the case of simple subprograms, both of these parts have fixed sizes.

The format, or layout, of the noncode part of a subprogram is called an activation record, because the data it describes are relevant only during the activation or execution of the subprogram. The form of an activation record is static. An activation record instance is a concrete example of an activation record, a collection of data in the form of an activation record.

Because languages with simple subprograms do not support recursion, there can be only one active version of a given subprogram at a time. Therefore, there can be only a single instance of the activation record for a subprogram. One possible layout for activation records is shown in Figure 10.1. The saved execution status of the caller is omitted here and in the remainder of this chapter because it is simple and not relevant to the discussion.

Because an activation record instance for a “simple” subprogram has fixed size, it can be statically allocated. In fact, it could be attached to the code part of the subprogram.

Figure 10.2 shows a program consisting of a main program and three subprograms: A, B, and C. Although the figure shows all the code segments separated from all the activation record instances, in some cases, the activation record instances are attached to their associated code segments.

The construction of the complete program shown in Figure 10.2 is not done entirely by the compiler. In fact, if the language allows independent compilation, the four program units—MAIN, A, B, and C—may have been compiled on different days, or even in different years. At the time each unit is compiled, the machine code for it, along with a list of references to external subprograms, is written to a file. The executable program shown in Figure 10.2 is put together by the linker, which is part of the operating system. (Sometimes linkers are called loaders, linker/loaders, or link editors.) When the linker is called for a main program, its first task is to find the files that contain the translated subprograms referenced in that program and load them into memory. Then, the linker must set the target addresses of all calls to those subprograms in the main program to the entry addresses of those subprograms. The same must be done for all calls to subprograms in the loaded subprograms and all calls to library subprograms. In the previous example, the linker was called for MAIN. The linker had to find the machine code programs for A, B, and C, along with their activation record instances, and load them into memory with the code for MAIN. Then, it had to
Implementing Subprograms with Stack-Dynamic Local Variables

We now examine the implementation of the subprogram linkage in languages in which locals are stack dynamic, again focusing on the call and return operations.

One of the most important advantages of stack-dynamic local variables is support for recursion. Therefore, languages that use stack-dynamic local variables also support recursion.

A discussion of the additional complexity required when subprograms can be nested is postponed until Section 10.4.

10.3.1 More Complex Activation Records

Subprogram linkage in languages that use stack-dynamic local variables are more complex than the linkage of simple subprograms for the following reasons:

- The compiler must generate code to cause the implicit allocation and deallocation of local variables.
• Recursion adds the possibility of multiple simultaneous activations of a subprogram, which means that there can be more than one instance (incomplete execution) of a subprogram at a given time, with at least one call from outside the subprogram and one or more recursive calls. The number of activations is limited only by the memory size of the machine. Each activation requires its activation record instance.

The format of an activation record for a given subprogram in most languages is known at compile time. In many cases, the size is also known for activation records because all local data are of a fixed size. That is not the case in some other languages, such as Ada, in which the size of a local array can depend on the value of an actual parameter. In those cases, the format is static, but the size can be dynamic. In languages with stack-dynamic local variables, activation record instances must be created dynamically. The typical activation record for such a language is shown in Figure 10.3.

Because the return address, dynamic link, and parameters are placed in the activation record instance by the caller, these entries must appear first.

The return address usually consists of a pointer to the instruction following the call in the code segment of the calling program unit. The dynamic link is a pointer to the base of the activation record instance of the caller. In static-scoped languages, this link is used to provide traceback information when a run-time error occurs. In dynamic-scoped languages, the dynamic link is used to access nonlocal variables. The actual parameters in the activation record are the values or addresses provided by the caller.

Local scalar variables are bound to storage within an activation record instance. Local variables that are structures are sometimes allocated elsewhere, and only their descriptors and a pointer to that storage are part of the activation record. Local variables are allocated and possibly initialized in the called subprogram, so they appear last.

Consider the following skeletal C function:

```c
void sub(float total, int part) {
    int list[5];
    float sum;
    . . .
}
```

**Figure 10.3**
A typical activation record for a language with stack-dynamic local variables
The activation record for `sub` is shown in Figure 10.4.

Activating a subprogram requires the dynamic creation of an instance of the activation record for the subprogram. As stated earlier, the format of the activation record is fixed at compile time, although its size may depend on the call in some languages. Because the call and return semantics specify that the subprogram last called is the first to complete, it is reasonable to create instances of these activation records on a stack. This stack is part of the run-time system and therefore is called the run-time stack, although we will usually just refer to it as the stack. Every subprogram activation, whether recursive or nonrecursive, creates a new instance of an activation record on the stack. This provides the required separate copies of the parameters, local variables, and return address.

One more thing is required to control the execution of a subprogram—the EP. Initially, the EP points at the base, or first address of the activation record instance of the main program. The run-time system must ensure that it always points at the base of the activation record instance of the currently executing program unit. When a subprogram is called, the current EP is saved in the new activation record instance as the dynamic link. The EP is then set to point at the base of the new activation record instance. Upon return from the subprogram, the stack top is set to the value of the current EP minus one and the EP is set to the dynamic link from the activation record instance of the subprogram that has completed its execution. Resetting the stack top effectively removes the top activation record instance.
The EP is used as the base of the offset addressing of the data contents of the activation record instance—parameters and local variables.

Note that the EP currently being used is not stored in the run-time stack. Only saved versions are stored in the activation record instances as the dynamic links.

We have now discussed several new actions in the linkage process. The lists given in Section 10.2 must be revised to take these into account. Using the activation record form given in this section, the new actions are as follows:

The caller actions are as follows:

1. Create an activation record instance.
2. Save the execution status of the current program unit.
3. Compute and pass the parameters.
4. Pass the return address to the called.
5. Transfer control to the called.

The prologue actions of the called are as follows:

1. Save the old EP in the stack as the dynamic link and create the new value.
2. Allocate local variables.

The epilogue actions of the called are as follows:

1. If there are pass-by-value-result or out-mode parameters, the current values of those parameters are moved to the corresponding actual parameters.
2. If the subprogram is a function, the functional value is moved to a place accessible to the caller.
3. Restore the stack pointer by setting it to the value of the current EP minus one and set the EP to the old dynamic link.
4. Restore the execution status of the caller.
5. Transfer control back to the caller.

Recall from Chapter 9, that a subprogram is active from the time it is called until the time that execution is completed. At the time it becomes inactive, its local scope ceases to exist and its referencing environment is no longer meaningful. Therefore, at that time, its activation record instance can be destroyed.

Parameters are not always transferred in the stack. In many compilers for RISC machines, parameters are passed in registers. This is because RISC machines normally have many more registers than CISC machines. In the remainder of this chapter, however, we assume that parameters are passed in
the stack. It is straightforward to modify this approach for parameters being passed in registers.

### 10.3.2 An Example Without Recursion

Consider the following skeletal C program:

```c
void fun1(float r) {
    int s, t;
    . . . 1
    fun2(s);
    . . .
}

void fun2(int x) {
    int y;
    . . . 2
    fun3(y);
    . .
}

void fun3(int q) {
    . . . 3
}

void main() {
    float p;
    . . .
    fun1(p);
    . . .
}
```

The sequence of function calls in this program is

main calls fun1
fun1 calls fun2
fun2 calls fun3

The stack contents for the points labeled 1, 2, and 3 are shown in Figure 10.5.

At point 1, only the activation record instances for function main and function fun1 are on the stack. When fun1 calls fun2, an instance of fun2’s activation record is created on the stack. When fun2 calls fun3, an instance of fun3’s activation record is created on the stack. When fun3's execution ends, the instance of its activation record is removed from the stack, and the EP is used to reset the stack top pointer. Similar processes take place when functions fun2 and fun1 terminate. After the return from the call to fun1 from main, the stack has only the instance of the activation record of main. Note that some implementations do not actually use an activation record instance on the stack.
for main functions, such as the one shown in the figure. However, it can be done this way, and it simplifies both the implementation and our discussion. In this example and in all others in this chapter, we assume that the stack grows from lower addresses to higher addresses, although in a particular implementation, the stack may grow in the opposite direction.

The collection of dynamic links present in the stack at a given time is called the **dynamic chain**, or **call chain**. It represents the dynamic history of how execution got to its current position, which is always in the subprogram code whose activation record instance is on top of the stack. References to local variables can be represented in the code as offsets from the beginning of the activation record of the local scope, whose address is stored in the EP. Such an offset is called a **local offset**.

The local_offset of a variable in an activation record can be determined at compile time, using the order, types, and sizes of variables declared in the subprogram associated with the activation record. To simplify the discussion, we assume that all variables take one position in the activation record. The first local variable declared in a subprogram would be allocated in the activation record two positions plus the number of parameters from the bottom (the first two positions are for the return address and the dynamic link). The second local...
10.3 Implementing Subprograms with Stack-Dynamic Local Variables

variable declared would be one position nearer the stack top and so forth. For example, consider the preceding example program. In fun1, the local_offset of s is 3; for t it is 4. Likewise, in fun2, the local_offset of y is 3. To get the address of any local variable, the local_offset of the variable is added to the EP.

10.3.3 Recursion

Consider the following example C program, which uses recursion to compute the factorial function:

```c
int factorial(int n) {
    if (n <= 1)
        return 1;
    else return (n * factorial(n - 1));
}

void main() {
    int value;
    value = factorial(3);
}
```

The activation record format for the function `factorial` is shown in Figure 10.6. Notice that it has an additional entry for the return value of the function.

Figure 10.7 shows the contents of the stack for the three times execution reaches position 1 in the function `factorial`. Each shows one more activation of the function, with its functional value undefined. The first activation record instance has the return address to the calling function, `main`. The others have a return address to the function itself; these are for the recursive calls.

Figure 10.8 shows the stack contents for the three times that execution reaches position 2 in the function `factorial`. Position 2 is meant to be the time after the `return` is executed but before the activation record has been removed from the stack. Recall that the code for the function multiplies the current value of the parameter `n` by the value returned by the recursive call to the function.

![Figure 10.6](image-url)

The activation record for `factorial`
The first return from `factorial` returns the value 1. The activation record instance for that activation has a value of 1 for its version of the parameter `n`. The result from that multiplication, 1, is returned to the second activation of `factorial` to be multiplied by its parameter value for `n`, which is 2. This step returns the value 2 to the first activation of `factorial` to be multiplied by its parameter value for `n`, which is 3, yielding the final functional value of 6, which is then returned to the first call to `factorial` in `main`. 

ARI = activation record instance

**Figure 10.7**

Stack contents at position 1 in `factorial`
Some of the non–C-based static-scoped programming languages use stack-dynamic local variables and allow subprograms to be nested. Among these are Fortran 95+, Ada, Python, JavaScript, Ruby, and Lua, as well as the functional languages. In this section, we examine the most commonly used approach to
implementing subprograms that may be nested. Until the very end of this section, we ignore closures.

10.4.1 The Basics

A reference to a nonlocal variable in a static-scoped language with nested subprograms requires a two-step access process. All nonstatic variables that can be nonlocally accessed are in existing activation record instances and therefore are somewhere in the stack. The first step of the access process is to find the instance of the activation record in the stack in which the variable was allocated. The second part is to use the local_offset of the variable (within the activation record instance) to access it.

Finding the correct activation record instance is the more interesting and more difficult of the two steps. First, note that in a given subprogram, only variables that are declared in static ancestor scopes are visible and can be accessed. Also, activation record instances of all of the static ancestors are always on the stack when variables in them are referenced by a nested subprogram. This is guaranteed by the static semantic rules of the static-scoped languages: A subprogram is callable only when all of its static ancestor subprograms are active.\(^1\) If a particular static ancestor were not active, its local variables would not be bound to storage, so it would be nonsense to allow access to them.

The semantics of nonlocal references dictates that the correct declaration is the first one found when looking through the enclosing scopes, most closely nested first. So, to support nonlocal references, it must be possible to find all of the instances of activation records in the stack that correspond to those static ancestors. This observation leads to the implementation approach described in the following subsection.

We do not address the issue of blocks until Section 10.5, so in the remainder of this section, all scopes are assumed to be defined by subprograms. Because functions cannot be nested in the C-based languages (the only static scope in those languages are those created with blocks), the discussions of this section do not apply to those languages directly.

10.4.2 Static Chains

The most common way to implement static scoping in languages that allow nested subprograms is static chaining. In this approach, a new pointer, called a static link, is added to the activation record. The **static link**, which is sometimes called a *static scope pointer*, points to the bottom of the activation record instance of an activation of the static parent. It is used for accesses to nonlocal variables. Typically, the static link appears in the activation record below the parameters. The addition of the static link to the activation record requires that local offsets differ from when the static link is not included. Instead of having two activation record elements before the parameters, there are now three: the return address, the static link, and the dynamic link.

---

1. Closures, of course, violate this rule.
A **static chain** is a chain of static links that connect certain activation record instances in the stack. During the execution of a subprogram \( P \), the static link of its activation record instance points to an activation record instance of \( P \)'s static parent program unit. That instance’s static link points in turn to \( P \)'s static grandparent program unit’s activation record instance, if there is one. So, the static chain connects all the static ancestors of an executing subprogram, in order of static parent first. This chain can obviously be used to implement the accesses to nonlocal variables in static-scoped languages.

Finding the correct activation record instance of a nonlocal variable using static links is relatively straightforward. When a reference is made to a nonlocal variable, the activation record instance containing the variable can be found by searching the static chain until a static ancestor activation record instance is found that contains the variable. However, it can be much easier than that. Because the nesting of scopes is known at compile time, the compiler can determine not only that a reference is nonlocal but also the length of the static chain that must be followed to reach the activation record instance that contains the nonlocal object.

Let **static_depth** be an integer associated with a static scope that indicates how deeply it is nested in the outermost scope. A program unit that is not nested inside any other unit has a static_depth of 0. If subprogram \( A \) is defined in a nonnested program unit, its static_depth is 1. If subprogram \( A \) contains the definition of a nested subprogram \( B \), then \( B \)'s static_depth is 2.

The length of the static chain needed to reach the correct activation record instance for a nonlocal reference to a variable \( x \) is exactly the difference between the static_depth of the subprogram containing the reference to \( x \) and the static_depth of the subprogram containing the declaration for \( x \). This difference is called the **nesting_depth**, or **chain_offset**, of the reference. The actual reference can be represented by an ordered pair of integers (chain_offset, local_offset), where chain_offset is the number of links to the correct activation record instance (local_offset is described in Section 10.3.2). For example, consider the following skeletal Python program:

```python
# Global scope
.
.
def f1():
    def f2():
        def f3():
            .
            # end of f3
            .
            # end of f2
        .
    # end of f1
```

The static_depths of the global scope, \( f1 \), \( f2 \), and \( f3 \) are 0, 1, 2, and 3, respectively. If procedure \( f3 \) references a variable declared in \( f1 \), the chain_offset of that reference would be 2 (static_depth of \( f3 \) minus the static_depth of \( f1 \)). If procedure \( f3 \) references a variable declared in \( f2 \), the chain_offset of that reference would be 1. References to locals can be handled using the same mechanism, with a chain_offset of 0, but instead of using the static pointer to the
activation record instance of the subprogram where the variable was declared as the base address, the EP is used.

To illustrate the complete process of nonlocal accesses, consider the following skeletal JavaScript program:

```javascript
function main() {
    var x;
    function bigsub() {
        var a, b, c;
        function sub1 {
            var a, d;
            ...
            a = b + c; <-----------------------------1
            ...
        } // end of sub1
        function sub2(x) {
            var b, e;
            function sub3() {
                var c, e;
                ...
                sub1();
                ...
                e = b + a; <-----------------------------2
            } // end of sub3
            ...
            sub3();
            ...
            a = d + e; <-----------------------------3
        } // end of sub2
        ...
        sub2(7);
        ...
    } // end of bigsub
    ...
    bigsub();
    ...
} // end of main
```

The sequence of procedure calls is

- `main` calls `bigsub`
- `bigsub` calls `sub2`
- `sub2` calls `sub3`
- `sub3` calls `sub1`

The stack situation when execution first arrives at point 1 in this program is shown in Figure 10.9.

At position 1 in procedure `sub1`, the reference is to the local variable, `a`, not to the nonlocal variable `a` from `bigsub`. This reference to `a` has the chain_offset/local_offset pair (0, 3). The reference to `b` is to the nonlocal `b` from `bigsub`. It can be represented by the pair (1, 4). The local_offset is 4, because a 3 offset
would be the first local variable (bigsub has no parameters). Notice that if the
dynamic link were used to do a simple search for an activation record instance
with a declaration for the variable b, it would find the variable b declared in
sub2, which would be incorrect. If the (1, 4) pair were used with the dynamic
chain, the variable e from sub3 would be used. The static link, however, points
to the activation record for bigsub, which has the correct version of b. The
variable b in sub2 is not in the referencing environment at this point and is
(correctly) not accessible. The reference to c at point 1 is to the c defined in
bigsub, which is represented by the pair (1, 5).

After sub1 completes its execution, the activation record instance for sub1
is removed from the stack, and control returns to sub3. The reference to the
variable e at position 2 in sub3 is local and uses the pair (0, 4) for access. The

Figure 10.9
Stack contents at position 1 in the program main

ARI = activation record instance
reference to the variable \( b \) is to the one declared in \( \text{sub2} \), because that is the nearest static ancestor that contains such a declaration. It is accessed with the pair \((1, 4)\). The local_offset is 4 because \( b \) is the first variable declared in \( \text{sub1} \), and \( \text{sub2} \) has one parameter. The reference to the variable \( a \) is to the \( a \) declared in \( \text{bigsub} \), because neither \( \text{sub3} \) nor its static parent \( \text{sub2} \) has a declaration for a variable named \( a \). It is referenced with the pair \((2, 3)\).

After \( \text{sub3} \) completes its execution, the activation record instance for \( \text{sub3} \) is removed from the stack, leaving only the activation record instances for \( \text{main} \), \( \text{bigsub} \), and \( \text{sub2} \). At position 3 in \( \text{sub2} \), the reference to the variable \( a \) is to the \( a \) in \( \text{bigsub} \), which has the only declaration of \( a \) among the active routines. This access is made with the pair \((1, 3)\). At this position, there is no visible scope containing a declaration for the variable \( d \), so this reference to \( d \) is a static semantics error. The error would be detected when the compiler attempted to compute the chain_offset/local_offset pair. The reference to \( e \) is to the local \( e \) in \( \text{sub2} \), which can be accessed with the pair \((0, 5)\).

In summary, the references to the variable \( a \) at points 1, 2, and 3 would be represented by the following points:

- \((0, 3)\) (local)
- \((2, 3)\) (two levels away)
- \((1, 3)\) (one level away)

It is reasonable at this point to ask how the static chain is maintained during program execution. If its maintenance is too complex, the fact that it is simple and effective would be unimportant. We assume here that parameters that are subprograms are not implemented.

The static chain must be modified for each subprogram call and return. The return part is trivial: When the subprogram terminates, its activation record instance is removed from the stack. After this removal, the new top activation record instance is that of the unit that called the subprogram whose execution just terminated. Because the static chain from this activation record instance was never changed, it works correctly just as it did before the call to the other subprogram. Therefore, no other action is required.

The action required at a subprogram call is more complex. Although the correct parent scope is easily determined at compile time, the most recent activation record instance of the parent scope must be found at the time of the call. This can be done by looking at activation record instances on the dynamic chain until the first one of the parent scope is found. However, this search can be avoided by treating subprogram declarations and references exactly like variable declarations and references. When the compiler encounters a subprogram call, among other things, it determines the subprogram that declared the called subprogram, which must be a static ancestor of the calling routine. It then computes the nesting_depth, or number of enclosing scopes between the caller and the subprogram that declared the called subprogram. This information is stored and can be accessed by the subprogram call during execution. At the time of the call, the static link of the called subprogram’s activation record instance is found by moving down the static chain of the caller. The number
of links in this move is equal to the nesting_depth, which was computed at compile time.

Consider again the program `main` and the stack situation shown in Figure 10.9. At the call to `sub1` in `sub3`, the compiler determines the nesting_depth of `sub3` (the caller) to be two levels inside the procedure that declared the called procedure `sub1`, which is `bigsub`. When the call to `sub1` in `sub3` is executed, this information is used to set the static link of the activation record instance for `sub1`. This static link is set to point to the activation record instance that is pointed to by the second static link in the static chain from the caller’s activation record instance. In this case, the caller is `sub3`, whose static link points to its parent’s activation record instance (that of `sub2`). The static link of the activation record instance for `sub2` points to the activation record instance for `bigsub`. So, the static link for the new activation record instance for `sub1` is set to point to the activation record instance for `bigsub`.

This method works for all subprogram linkage, except when parameters that are subprograms are involved.

One criticism of using the static chain approach to access nonlocal variables is that references to variables in scopes beyond the static parent cost more than references to locals. The static chain must be followed, one link per enclosing scope from the reference to the declaration. Fortunately, in practice, references to distant nonlocal variables are rare, so this is not a serious problem. Another criticism of the static-chain approach is that it is difficult for a programmer working on a time-critical program to estimate the costs of nonlocal references, because the cost of each reference depends on the depth of nesting between the reference and the scope of declaration. Further complicating this problem is that subsequent code modifications may change nesting depths, thereby changing the timing of some references, both in the changed code and possibly in code far from the changes.

Some alternatives to static chains have been developed, most notably an approach that uses an auxiliary data structure called a display. However, none of the alternatives has been found to be superior to the static-chain method, which is still the most widely used approach. Therefore, none of the alternatives are discussed here.

The processes and data structures described in this section correctly implement closures in languages that do not permit functions to return functions and do not allow functions to be assigned to variables. However, they are inadequate for languages that do allow one or both of those operations. Several new mechanisms are needed to implement access to nonlocals in such languages. First, if a subprogram accesses a variable from a nesting but not global scope, that variable cannot be stored only in the activation record of its home scope. That activation record could be deallocated before the subprogram that needs it is activated. Such variables could also be stored in the heap and given unlimited extend (their lifetimes are the lifetime of the whole program). Second, subprograms must have mechanisms to access the nonlocals that are stored in the heap. Third, the heap-allocated variables that are nonlocally accessed must be updated every time their stack versions are updated. Clearly, these are nontrivial extensions to the implementation static scoping using static chains.
Recall from Chapter 5, that a number of languages, including the C-based languages, provide for user-specified local scopes for variables called **blocks**. As an example of a block, consider the following code segment:

```c
{ int temp;
  temp = list[upper];
  list[upper] = list[lower];
  list[lower] = temp;
}
```

A block is specified in the C-based languages as a compound statement that begins with one or more data definitions. The lifetime of the variable `temp` in the preceding block begins when control enters the block and ends when control exits the block. The advantage of using such a local is that it cannot interfere with any other variable with the same name that is declared elsewhere in the program, or more specifically, in the referencing environment of the block.

Blocks can be implemented by using the static-chain process described in Section 10.4 for implementing nested subprograms. Blocks are treated as parameterless subprograms that are always called from the same place in the program. Therefore, every block has an activation record. An instance of its activation record is created every time the block is executed.

Blocks can also be implemented in a different and somewhat simpler and more efficient way. The maximum amount of storage required for block variables at any time during the execution of a program can be statically determined, because blocks are entered and exited in strictly textual order. This amount of space can be allocated after the local variables in the activation record. Offsets for all block variables can be statically computed, so block variables can be addressed exactly as if they were local variables.

For example, consider the following skeletal program:

```c
void main() {
  int x, y, z;
  while ( . . ) {
    int a, b, c;
    .
    while ( . . . ) {
      int d, e;
      .
      }
    }
  while ( . . . ) {
    int f, g;
    . .
    }
  . .
}
```
For this program, the static-memory layout shown in Figure 10.10 could be used. Note that $f$ and $g$ occupy the same memory locations as $a$ and $b$, because $a$ and $b$ are popped off the stack when their block is exited (before $f$ and $g$ are allocated).

### 10.6 Implementing Dynamic Scoping

There are at least two distinct ways in which local variables and nonlocal references to them can be implemented in a dynamic-scoped language: deep access and shallow access. Note that deep access and shallow access are not concepts related to deep and shallow binding. An important difference between binding and access is that deep and shallow bindings result in different semantics; deep and shallow accesses do not.

#### 10.6.1 Deep Access

If local variables are stack dynamic and are part of the activation records in a dynamic-scoped language, references to nonlocal variables can be resolved by searching through the activation record instances of the other subprograms.
that are currently active, beginning with the one most recently activated. This concept is similar to that of accessing nonlocal variables in a static-scoped language with nested subprograms, except that the dynamic—rather than the static—chain is followed. The dynamic chain links together all subprogram activation record instances in the reverse of the order in which they were activated. Therefore, the dynamic chain is exactly what is needed to reference nonlocal variables in a dynamic-scoped language. This method is called **deep access**, because access may require searches deep into the stack.

Consider the following example skeletal program:

```c
void sub3() {
    int x, z;
    x = u + v;
    . . .
}

void sub2() {
    int w, x;
    . . .
}

void sub1() {
    int v, w;
    . . .
}

void main() {
    int v, u;
    . . .
}
```

This program is written in a syntax that gives it the appearance of a program in a C-based language, but it is not meant to be in any particular language. Suppose the following sequence of function calls occurs:

- `main` calls `sub1`
- `sub1` calls `sub1`
- `sub1` calls `sub2`
- `sub2` calls `sub3`

Figure 10.11 shows the stack during the execution of function `sub3` after this calling sequence. Notice that the activation record instances do not have static links, which would serve no purpose in a dynamic-scoped language.

Consider the references to the variables `x`, `u`, and `v` in function `sub3`. The reference to `x` is found in the activation record instance for `sub3`. The reference to `u` is found by searching all of the activation record instances on the stack, because the only existing variable with that name is in `main`. This search involves following four dynamic links and examining 10 variable names. The reference to `v` is found in the most recent (nearest on the dynamic chain) activation record instance for the subprogram `sub1`. 
There are two important differences between the deep-access method for nonlocal access in a dynamic-scoped language and the static-chain method for static-scoped languages. First, in a dynamic-scoped language, there is no way to determine at compile time the length of the chain that must be searched. Every activation record instance in the chain must be searched until the first instance of the variable is found. This is one reason why dynamic-scoped languages typically have slower execution speeds than static-scoped languages. Second, activation records must store the names of variables for the search process, whereas in static-scoped language implementations only the values are required. (Names are not required for static scoping, because all variables are represented by the chain_offset/local_offset pairs.)

### 10.6.2 Shallow Access

Shallow access is an alternative implementation method, not an alternative semantics. As stated previously, the semantics of deep access and shallow access are identical. In the shallow-access method, variables declared in subprograms are not stored in the activation records of those subprograms. Because with dynamic scoping there is at most one visible version of a variable of any specific name at a given time, a very different approach can be taken. One variation of
Implementing Subprograms

shallow access is to have a separate stack for each variable name in a complete program. Every time a new variable with a particular name is created by a declaration at the beginning of a subprogram that has been called, the variable is given a cell at the top of the stack for its name. Every reference to the name is to the variable on top of the stack associated with that name, because the top one is the most recently created. When a subprogram terminates, the lifetimes of its local variables end, and the stacks for those variable names are popped. This method allows fast references to variables, but maintaining the stacks at the entrances and exits of subprograms is costly.

Figure 10.12 shows the variable stacks for the earlier example program in the same situation as shown with the stack in Figure 10.11.

Another option for implementing shallow access is to use a central table that has a location for each different variable name in a program. Along with each entry, a bit called active is maintained that indicates whether the name has a current binding or variable association. Any access to any variable can then be to an offset into the central table. The offset is static, so the access can be fast. SNOBOL implementations use the central table implementation technique.

Maintenance of a central table is straightforward. A subprogram call requires that all of its local variables be logically placed in the central table. If the position of the new variable in the central table is already active—that is, if it contains a variable whose lifetime has not yet ended (which is indicated by the active bit)—that value must be saved somewhere during the lifetime of the new variable. Whenever a variable begins its lifetime, the active bit in its central table position must be set.

There have been several variations in the design of the central table and in the way values are stored when they are temporarily replaced. One variation is to have a “hidden” stack on which all saved objects are stored. Because subprogram calls and returns, and thus the lifetimes of local variables, are nested, this works well.

The second variation is perhaps the cleanest and least expensive to implement. A central table of single cells is used, storing only the current version of each variable with a unique name. Replaced variables are stored in the activation record of the subprogram that created the replacement variable. This is a stack mechanism, but it uses the stack that already exists, so the new overhead is minimal.

The choice between shallow and deep access to nonlocal variables depends on the relative frequencies of subprogram calls and nonlocal references. The deep-access method provides fast subprogram linkage, but references to nonlocals,

---

**Figure 10.12**

One method of using shallow access to implement dynamic scoping

(The names in the stack cells indicate the program units of the variable declaration.)
especially references to distant nonlocals (in terms of the call chain), are costly. The shallow-access method provides much faster references to nonlocals, especially distant nonlocals, but is more costly in terms of subprogram linkage.

**Summary**

Subprogram linkage semantics requires many actions by the implementation. In the case of “simple” subprograms, these actions are relatively uncomplicated. At the call, the status of execution must be saved, parameters and the return address must be passed to the called subprogram, and control must be transferred. At the return, the values of pass-by-result and pass-by-value-result parameters must be transferred back, as well as the return value if it is a function, execution status must be restored, and control transferred back to the caller. In languages with stack-dynamic local variables and nested subprograms, subprogram linkage is more complex. There may be more than one activation record instance, those instances must be stored on the run-time stack, and static and dynamic links must be maintained in the activation record instances. The static link is to allow references to nonlocal variables in static-scoped languages.

Subprograms in languages with stack-dynamic local variables and nested subprograms have two components: the actual code, which is static, and the activation record, which is stack dynamic. Activation record instances contain the formal parameters and local variables, among other things.

Access to nonlocal variables in a dynamic-scoped language can be implemented by use of the dynamic chain or through some central variable table method. Dynamic chains provide slow accesses but fast calls and returns. The central table methods provide fast accesses but slow calls and returns.

**Review Questions**

1. What is the definition used in this chapter for “simple” subprograms?
2. Which of the caller or callee saves execution status information?
3. What must be stored for the linkage to a subprogram?
4. What is the task of a linker?
5. What are the two reasons why implementing subprograms with stack-dynamic local variables is more difficult than implementing simple subprograms?
6. What is the difference between an activation record and an activation record instance?
7. Why are the return address, dynamic link, and parameters placed in the bottom of the activation record?
8. What kind of machines often use registers to pass parameters?
9. What are the two steps in locating a nonlocal variable in a static-scoped language with stack-dynamic local variables and nested subprograms?

10. Define static chain, static_depth, nesting_depth, and chain_offset.

11. What is an EP, and what is its purpose?

12. How are references to variables represented in the static-chain method?

13. Name three widely used programming languages that do not allow nested subprograms.

14. What are the two potential problems with the static-chain method?

15. Explain the two methods of implementing blocks.

16. Describe the deep-access method of implementing dynamic scoping.

17. Describe the shallow-access method of implementing dynamic scoping.

18. What are the two differences between the deep-access method for nonlocal access in dynamic-scoped languages and the static-chain method for static-scoped languages?

19. Compare the efficiency of the deep-access method to that of the shallow-access method, in terms of both calls and nonlocal accesses.

**Problem Set**

1. Show the stack with all activation record instances, including static and dynamic chains, when execution reaches position 1 in the following skeletal program. Assume bigsub is at level 1.

   ```
   function bigsub() {
     function a() {
       function b() {
         ...
         // end of b
     function c() {
       ...
       b();
       ...
       // end of c
       ...
       c();
       ...
     } // end of a
     ...
     a();
     ...
   } // end of bigsub
   ```
2. Show the stack with all activation record instances, including static and dynamic chains, when execution reaches position 1 in the following skeletal program. Assume bigsub is at level 1.

```javascript
function bigsub() {
    var mysum;
    function a() {
        var x;
        function b(sum) {
            var y, z;
            ...
            c(z);
            ...
        } // end of b
        ...
        b(x);
        ...
    } // end of a
    function c(plums) {
        ...
    } // end of c
    var l;
    ...
    a();
    ...
} // end of bigsub
```

3. Show the stack with all activation record instances, including static and dynamic chains, when execution reaches position 1 in the following skeletal program. Assume bigsub is at level 1.

```javascript
function bigsub() {
    function a(flag) {
        function b() {
            ...
            a(false);
            ...
        } // end of b
        ...
        if (flag)
            b();
        else c();
        ...
    } // end of a
    function c() {
        function d() {
            ...
        } // end of d
        ...
    d();
```
4. Show the stack with all activation record instances, including static and dynamic chains, when execution reaches position 1 in the following skeletal program. This program uses the deep-access method to implement dynamic scoping.

```c
void fun1() {
    float a;
    . . .
}

void fun2() {
    int b, c;
    . . . <----------- 1
}

void fun3() {
    float d;
    . . .
}

void main() {
    char e, f, g;
    . . .
}
```

The calling sequence for this program for execution to reach `fun3` is

- `main` calls `fun1`
- `fun1` calls `fun3`
- `fun3` calls `fun2`

5. Assume that the program of Problem 4 is implemented using the shallow-access method using a stack for each variable name. Show the stacks for the time of the execution of `fun3`, assuming execution found its way to that point through the sequence of calls shown in Problem 4.
6. Although local variables in the methods of C-based languages are allocated at the beginning of each activation, under what circumstances could the value of a local variable in a particular activation retain the value of the previous activation?

7. It is stated in this chapter that when nonlocal variables are accessed in a dynamic-scoped language using the dynamic chain, variable names must be stored in the activation records with the values. If this were actually done, every nonlocal access would require a sequence of costly string comparisons on names. Design an alternative to these string comparisons that would be faster.

8. Pascal allows gotos with nonlocal targets. How could such statements be handled if static chains were used for nonlocal variable access? *Hint:* Consider the way the correct activation record instance of the static parent of a newly enacted procedure is found (see Section 10.4.2).

9. The static-chain method could be expanded slightly by using two static links in each activation record instance where the second points to the static grandparent activation record instance. How would this approach affect the time required for subprogram linkage and nonlocal references?

10. Design a skeletal program and a calling sequence that results in an activation record instance in which the static and dynamic links point to different activation-recorded instances in the run-time stack.

11. If a compiler uses the static chain approach to implementing blocks, which of the entries in the activation records for subprograms are needed in the activation records for blocks?

12. Examine the subprogram call instructions of three different architectures, including at least one CISC machine and one RISC machine, and write a short comparison of their capabilities. (The design of these instructions usually determines at least part of the compiler writer’s design of subprogram linkage.)

---

**Programming Exercises**

1. Write a program that includes two subprograms, one that takes a single parameter and performs some simple operation on that parameter and one that takes 20 parameters and uses all of the parameters, but only for one simple operation. The main program must call these two subprograms a large number of times. Include in the program timing code to output the run time of the calls to each of the two subprograms. Run the program on a RISC machine and on a CISC machine and compare the ratios of the time required by the two subprograms. Based on the results, what can you say about the speed of parameter passing on the two machines?
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Abstract Data Types and Encapsulation Constructs

11.1 The Concept of Abstraction
11.2 Introduction to Data Abstraction
11.3 Design Issues for Abstract Data Types
11.4 Language Examples
11.5 Parameterized Abstract Data Types
11.6 Encapsulation Constructs
11.7 Naming Encapsulations
In this chapter, we explore programming language constructs that support data abstraction. Among the new ideas of the last 50 years in programming methodologies and programming language design, data abstraction is one of the most profound.

We begin by discussing the general concept of abstraction in programming and programming languages. Data abstraction is then defined and illustrated with an example. This topic is followed by descriptions of the support for data abstraction in C++, Objective-C, Java, C#, and Ruby. To illuminate the similarities and differences in the design of the language facilities that support data abstraction, implementations of the same example data abstraction are given in C++, Objective-C, Java, and Ruby. Next, the capabilities of C++, Java 5.0, and C# 2005 to build parameterized abstract data types are discussed.

All the languages used in this chapter to illustrate the concepts and constructs of abstract data types support object-oriented programming, because virtually all contemporary languages support object-oriented programming and nearly all of those that do not, and yet support abstract data types, have faded into obscurity.

Constructs that support abstract data types are encapsulations of the data and operations on objects of the type. Encapsulations that contain multiple types are required for the construction of larger programs. These encapsulations and the associated namespace issues are also discussed in this chapter.

Some programming languages support logical, as opposed to physical, encapsulations, which are actually used to encapsulate names. These are discussed in Section 11.7.

11.1 The Concept of Abstraction

An abstraction is a view or representation of an entity that includes only the most significant attributes. In a general sense, abstraction allows one to collect instances of entities into groups in which their common attributes need not be considered. For example, suppose we define birds to be creatures with the following attributes: two wings, two legs, a tail, and feathers. Then, if we say a crow is a bird, a description of a crow need not include those attributes. The same is true for robins, sparrows, and yellow-bellied sapsuckers. The common attributes in the descriptions of specific species of birds can be abstracted away, because all species have them. Within a particular species, only the attributes that distinguish that species need be considered. For example, crows have the attributes of being black, being of a particular size, and being noisy. A description of a crow needs to provide those attributes, but not the others that are common to all birds. This results in significant simplification of the descriptions of members of the species. A less abstract view of a species, that of a bird, may be considered when it is necessary to see a higher level of detail, rather than just the special attributes.

In the world of programming languages, abstraction is a weapon against the complexity of programming; its purpose is to simplify the programming process. It is an effective weapon because it allows programmers to focus on essential attributes, while ignoring subordinate attributes.
The two fundamental kinds of abstraction in contemporary programming languages are process abstraction and data abstraction.

The concept of **process abstraction** is among the oldest in programming language design (Plankalkül supported process abstraction in the 1940s). All subprograms are process abstractions because they provide a way for a program to specify a process, without providing the details of how it performs its task (at least in the calling program). For example, when a program needs to sort an array of numeric data of some type, it usually uses a subprogram for the sorting process. At the point where the sorting process is required, a statement such as

```
sortInt(list, listLen)
```

is placed in the program. This call is an abstraction of the actual sorting process, whose algorithm is not specified. The call is independent of the algorithm implemented in the called subprogram.

In the case of the subprogram `sortInt`, the only essential attributes are the name of the array to be sorted, the type of its elements, the array's length, and the fact that the call to `sortInt` will result in the array being sorted. The particular algorithm that `sortInt` implements is an attribute that is not essential to the user. The user needs to see only the name and protocol of the sorting subprogram to be able to use it.

The widespread use of data abstraction necessarily followed that of process abstraction because an integral and essential part of every data abstraction is its operations, which are defined as process abstractions.

**11.2 Introduction to Data Abstraction**

The evolution of data abstraction began in 1960 with the first version of COBOL, which included the record data structure.1 The C-based languages have structs, which are also records. An abstract data type is a data structure, in the form of a record, but which includes subprograms that manipulate its data.

Syntactically, an abstract data type is an enclosure that includes only the data representation of one specific data type and the subprograms that provide the operations for that type. Through access controls, unnecessary details of the type can be hidden from units outside the enclosure that use the type. Program units that use an abstract data type can declare variables of that type, even though the actual representation is hidden from them. An instance of an abstract data type is called an **object**.

One of the motivations for data abstraction is similar to that of process abstraction. It is a weapon against complexity; a means of making large and/or complicated programs more manageable. Other motivations for and advantages of abstract data types are discussed later in this section.

---

1. Recall from Chapter 2 that a record is a data structure that stores fields, which have names and can be of different types.
Object-oriented programming, which is described in Chapter 12, is an outgrowth of the use of data abstraction in software development, and data abstraction is one of its fundamental components.

11.2.1 Floating-Point as an Abstract Data Type

The concept of an abstract data type, at least in terms of built-in types, is not a recent development. All built-in data types, even those of Fortran I, are abstract data types, although they are rarely called that. For example, consider a floating-point data type. Most programming languages include at least one of these. A floating-point type provides the means to create variables to store floating-point data and also provides a set of arithmetic operations for manipulating objects of the type.

Floating-point types in high-level languages employ a key concept in data abstraction: information hiding. The actual format of the floating-point data value in a memory cell is hidden from the user, and the only operations available are those provided by the language. The user is not allowed to create new operations on data of the type, except those that can be constructed using the built-in operations. The user cannot directly manipulate the parts of the actual representation of values because that representation is hidden. It is this feature that allows program portability between implementations of a particular language, even though the implementations may use different representations for particular data types. For example, before the IEEE 754 standard floating-point representations appeared in the mid-1980s, there were several different representations being used by different computer architectures. However, this variation did not prevent programs that used floating-point types from being portable among the various architectures.

11.2.2 User-Defined Abstract Data Types

A user-defined abstract data type should provide the same characteristics as those of language-defined types, such as a floating-point type: (1) a type definition that allows program units to declare variables of the type but hides the representation of objects of the type; and (2) a set of operations for manipulating objects of the type.

We now formally define an abstract data type in the context of user-defined types. An abstract data type is a data type that satisfies the following conditions:

- The representation of objects of the type is hidden from the program units that use the type, so the only direct operations possible on those objects are those provided in the type’s definition.
- The declarations of the type and the protocols of the operations on objects of the type, which provide the type’s interface, are contained in a single syntactic unit. The type’s interface does not depend on the representation of the objects or the implementation of the operations. Also, other program units are allowed to create variables of the defined type.

There are several benefits of information hiding. One of these is increased reliability. Program units that use a specific abstract data type are called clients of that type. Clients cannot manipulate the underlying representations of
objects directly, either intentionally or by accident, thus increasing the integrity of such objects. Objects can be changed only through the provided operations.

Another benefit of information hiding is it reduces the range of code and number of variables of which a programmer must be aware when writing or reading a part of the program. The value of a particular variable can be changed only by code in a restricted range, making the code easier to understand and making it less challenging to find sources of incorrect changes.

Information hiding also makes name conflicts less likely, because the scopes of variables is smaller.

Finally, consider the following advantage of information hiding: Suppose that the original implementation of the stack abstraction uses a linked list representation. At a later time, because of memory management problems with that representation, the stack abstraction is changed to use a contiguous representation (one that implements a stack in an array). Because data abstraction was used, this change can be made in the code that defines the stack type, but no changes will be required in any of the clients of the stack abstraction. Of course, a change in protocol of any of the operations would require changes in the clients.

Although the definition of an abstract data type specifies that data members of objects must be hidden from clients, many situations arise in which clients need to access these data members. The common solution is to provide accessor methods, sometimes called **getters** and **setters**, that allow clients indirect access to the so-called hidden data—a better solution than simply making the data public, which would provide direct access. There are three reasons why accessors are better:

1. Read-only access can be provided by having a getter method but no corresponding setter method.
2. Constraints can be included in setters. For example, if the data value should be restricted to a particular range, the setter can enforce that.
3. The actual implementation of the data member can be changed without affecting the clients if getters and setters are the only access.

Both specifying data in an abstract data type to be public and providing accessor methods for that data are violations of the principles of abstract data types. Some believe these are simply loopholes that make an imperfect design usable. As we will see in Section 11.4.5.2, Ruby disallows making instance data public. However, Ruby also makes it very easy to create accessor functions. It is a challenge for developers to design abstract data types in which all of the data is actually hidden.

The primary advantage of packaging the declarations of the type and its operations in a single syntactic unit is that it provides a method of organizing a program into logical units that can be compiled separately. In some cases, the implementation is included with the type declaration; in other cases, it is in a separate syntactic unit. The advantage of having the implementation of the type and its operations in different syntactic units is that it increases the program’s modularity and it is a clear separation of design and implementation. If both the declarations and the definitions of types and operations are in the same syntactic unit, there must be some means of hiding from client program units the parts of the unit that specify the definitions.
11.2.3 An Example

A stack is a widely applicable data structure that stores some number of data elements and only allows access to the data element at one of its ends, the top. Suppose an abstract data type is to be constructed for a stack that has the following abstract operations:

<table>
<thead>
<tr>
<th>Operation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>create(stack)</td>
<td>Creates and possibly initializes a stack object</td>
</tr>
<tr>
<td>destroy(stack)</td>
<td>Deallocates the storage for the stack</td>
</tr>
<tr>
<td>empty(stack)</td>
<td>A predicate (or Boolean) function that returns true if the specified stack is empty and false otherwise</td>
</tr>
<tr>
<td>push(stack, element)</td>
<td>Pushes the specified element on the specified stack</td>
</tr>
<tr>
<td>pop(stack)</td>
<td>Removes the top element from the specified stack</td>
</tr>
<tr>
<td>top(stack)</td>
<td>Returns a copy of the top element from the specified stack</td>
</tr>
</tbody>
</table>

Note that some implementations of abstract data types do not require the create and destroy operations. For example, simply defining a variable to be of an abstract data type may implicitly create the underlying data structure and initialize it. The storage for such a variable may be implicitly deallocated at the end of the variable's scope.

A client of the stack type could have a code sequence such as the following:

```plaintext
... create(stk1);
push(stk1, color1);
push(stk1, color2);
temp = top(stk1);
...
```

11.3 Design Issues for Abstract Data Types

A facility for defining abstract data types in a language must provide a syntactic unit that encloses the declaration of the type and the prototypes of the subprograms that implement the operations on objects of the type. It must be possible to make these visible to clients of the abstraction. This allows clients to declare variables of the abstract type and manipulate their values. Although the type name must have external visibility, the type representation must be hidden. The type representation and the definitions of the subprograms that implement the operations may appear inside or outside this syntactic unit.

Few, if any, general built-in operations should be provided for objects of abstract data types, other than those provided with the type definition. There simply are not many operations that apply to a broad range of abstract data
types. Among these are assignment and comparisons for equality and inequality. If the language does not allow users to overload assignment, the assignment operation must be included in the abstraction. Comparisons for equality and inequality should be predefined in the abstraction in some cases but not in others. For example, if the type is implemented as a pointer, equality may mean pointer equality, but the designer may want it to mean equality of the structures referenced by the pointers.

Some operations are required by many abstract data types, but because they are not universal, they often must be provided by the designer of the type. Among these are iterators, accessors, constructors, and destructors. Iterators were discussed in Chapter 8. Accessors provide a form of access to data that is hidden from direct access by clients. Constructors are used to initialize parts of newly created objects. Destructors are often used to reclaim heap storage that may be used by parts of abstract data type objects in languages that do not do implicit storage reclamation.

As stated earlier, the enclosure for an abstract data type defines a single data type and its operations. Many contemporary languages, including C++, Objective-C, Java, and C#, directly support abstract data types.

The first design issue is whether abstract data types can be parameterized. For example, if the language supports parameterized abstract data types, one could design an abstract data type for some structure that could store elements of any type. Parameterized abstract data types are discussed in Section 11.5. The second design issue is what access controls are provided and how such controls are specified. Finally, the language designer must decide whether the specification of the type is physically separate from its implementation (or whether that is a developer choice).

## 11.4 Language Examples

The concept of data abstraction had its origins in SIMULA 67, although that language did not provide complete support for abstract data types, because it did not include a way to hide implementation details. In this section, we describe the support for data abstraction provided by C++, Objective-C, Java, C#, and Ruby.

### 11.4.1 Abstract Data Types in C++

C++, which was first released in 1985, was created by adding features to C. The first important additions were those to support object-oriented programming. Because one of the primary components of object-oriented programming is abstract data types, C++ obviously is required to support them.

C++ provides two constructs that are very similar to each other, the class and the struct, which more directly support abstract data types. Because structs are most commonly used when only data is included, we do not discuss them further here.
C++: Its Birth, Its Ubiquitousness, and Common Criticisms

Bjarne Stroustrup

Bjarne Stroustrup is the designer and original implementer of C++ and the author of A Tour of C++, Programming – Principles and Practice using C++, The C++ Programming Language, The Design and Evolution of C++, and many other publications. His research interests include distributed systems, design, programming techniques, software development tools, and programming languages. He is actively involved in the ANSI/ISO standardization of C++. Dr. Stroustrup is a Managing Director in the technology division of Morgan Stanley in New York City, a Visiting Professor in Computer Science at Columbia University, and a Distinguished Research Professor in Computer Science at Texas A&M University. He is a member of the National Academy of Engineering, an ACM Fellow, and an IEEE fellow. In 1993, Stroustrup received the ACM Grace Murray Hopper Award “for his early work laying the foundations for the C++ programming language. Based on the foundations and Dr. Stroustrup’s continuing efforts, C++ has become one of the most influential programming languages in the history of computing.”

A BRIEF HISTORY OF YOU AND COMPUTING

What were you working on, and where, before you joined Bell Labs in the early 1980s? At Bell Labs, I was doing research in the general area of distributed systems. I joined in 1979. Before that, I was finishing my Ph.D. in that field in Cambridge University.

Did you immediately start on “C with Classes” (which would later become C++)? I worked on a few projects related to distributed computing before starting on C with Classes and during the development of that and of C++. For example, I was trying to find a way to distribute the UNIX kernel across several computers and helped a lot of projects build simulators.

Was it an interest in mathematics that got you into this profession? I signed up for a degree in “mathematics with computer science” and my master’s degree is officially a math degree. I—wrongly—thought that computing was some kind of applied math. I did a couple of years of math and rate myself a poor mathematician, but that’s still much better than not knowing math. At the time I signed up, I had never even seen a computer. What I love about computing is the programming rather than the more mathematical fields.

DISSECTING A SUCCESSFUL LANGUAGE

I’d like to work backward, listing some items I think make C++ ubiquitous, and get your reaction. It’s “open source,” nonproprietary, and standardized by ANSI/ISO. The ISO C++ standard is important. There are many independently developed and evolving C++ implementations. Without a standard for them to adhere to and a standards process to help coordinate the evolution of C++, a chaos of dialects would erupt.

It is also important that there are both open-source and commercial implementations available. In addition, for many users, it is crucial that the standard provides a measure of protection from manipulation by implementation providers.

The ISO standards process is open and democratic. The C++ committee rarely meets with fewer than 50 people present and typically more than eight nations are represented at each meeting. It is not just a vendors’ forum.
It’s ideal for systems programming (which, at the time C++ was born, was the largest sector of the market developing code).

Yes, C++ is a strong contender for any systems-programming project. It is also effective for embedded systems programming, which is currently the fastest-growing sector. Yet another growth area for C++ is high-performance numeric/engineering/scientific programming.

**Its object-oriented nature and inclusion of classes/libraries make programming more efficient and transparent.** C++ is a multiparadigm programming language. That is, it supports several fundamental styles of programming (including object-oriented programming) and combinations of those styles. When used well, this leads to cleaner, more flexible, and more efficient libraries than can be provided using just one paradigm. The C++ standard library containers and algorithms, which is basically a generic programming framework, is an example. When used together with (object-oriented) class hierarchies, the result is an unsurpassed combination of type safety, efficiency, and flexibility.

**Its incubation in the AT&T development environment** AT&T Bell Labs provided an environment that was crucial for C++’s development. The labs were an exceptionally rich source of challenging problems and a uniquely supportive environment for practical research. C++ emerged from the same research lab as C did and benefited from the same intellectual tradition, experience, and exceptional people. Throughout, AT&T supported the standardization of C++. However, C++ was not the beneficiary of a massive marketing campaign, like many modern languages. That’s simply not the way the labs work.

**Did I miss anything on your top list?** Undoubtedly.

**Now, let me paraphrase from the C++ critiques and get your reactions:** It’s huge/unwieldy. The “hello world” problem is 10 times larger in C++ than in C. C++ is certainly not a small language, but then few modern languages are. If a language is small, you tend to need huge libraries to get work done and often have to rely on conventions and extensions. I prefer to have key parts of the inevitable complexity in the language where it can be seen, taught, and effectively standardized rather than hidden elsewhere in a system. For most purposes, I don’t consider C++ unwieldy. The C++ “hello world” program isn’t larger than its C equivalent on my machine, and it shouldn’t be on yours.

In fact, the object code for the C++ version of the “hello world” program is smaller than the C version on my machine. There is no language reason why the one version should be larger than the other. It is all an issue of how the implementor organized the libraries. If one version is significantly larger than the other, report the problem to the implementor of the larger version.

**It’s tougher to program in C++ (compared with C).** (Something the critics say,) Even you once admitted it, saying something about shooting yourself in the foot with C versus C++. Yes, I did say something along the lines of “C makes it easy to shoot yourself in the foot; C++ makes it harder, but when you do, C++ blows your whole leg off.” What people tend to miss is that what I said about C++ is to a varying extent true for all powerful languages. As you protect people from simple dangers, they get themselves into new and less obvious problems. Someone who avoids the simple problems may simply be heading for a not-so-simple one. One problem with very supporting and protective environments is that the hard problems may be discovered too late or be too hard to remedy once discovered. Also, a rare problem is harder to find than a frequent one because you don’t suspect it.

**It’s appropriate for embedded systems of today but not for the Internet software of today.** C++ is suitable for embedded systems today. It is also suitable—and widely used—for “Internet software” today. For example, have a look at my “C++ applications” Web page. You’ll notice that some of the major Web service providers, such as Amazon, Adobe, Google, Quicken, and Microsoft, critically rely on C++. Gaming is a related area in which you find heavy C++ use.

**Did I miss another one that you get a lot?** Sure.
C++ classes are types. A C++ program unit that declares an instance of a class can also access any of the public entities in that class, but only through an instance of the class.

### 11.4.1.1 Encapsulation

The data defined in a C++ class are called **data members**; the functions (methods) defined in a class are called **member functions**. Data members and member functions appear in two categories: class and instance. Class members are associated with the class; instance members are associated with the instances of the class. In this chapter, only the instance members of a class are discussed. All of the instances of a class share a single set of member functions, but each instance has its own set of the class’s data members. Class instances can be static, stack dynamic, or heap dynamic. If static or stack dynamic, they are referenced directly with value variables. If heap dynamic, they are referenced through pointers. Stack dynamic instances of classes are always created by the elaboration of an object declaration. Furthermore, the lifetime of such a class instance ends when the end of the scope of its declaration is reached. Heap dynamic class instances are created with the `new` operator and destroyed with the `delete` operator. Both stack- and heap-dynamic classes can have pointer data members that reference heap dynamic data, so that even though a class instance is stack dynamic, it can include data members that reference heap dynamic data.

A member function of a class can be defined in two distinct ways: The complete definition can appear in the class, or only in its header. When both the header and the body of a member function appear in the class definition, the member function is implicitly inlined. Recall that this means that its code is placed in the caller’s code, rather than requiring the usual call and return linkage. If only the header of a member function appears in the class definition, its complete definition appears outside the class and is separately compiled. The rationale for allowing member functions to be inlined was to save function call overhead in real-time applications, in which run-time efficiency is of utmost importance. The downside of inlining member functions is that it clutters the class definition interface, resulting in a reduction in readability.

Placing member function definitions outside the class definition separates specification from implementation, a common goal of modern programming.

### 11.4.1.2 Information Hiding

A C++ class can contain both hidden and visible entities (meaning they are either hidden from or visible to clients of the class). Entities that are to be hidden are placed in a **private** clause, and visible, or public, entities appear in a **public** clause. The **public** clause therefore describes the interface to class instances. There is also a third category of visibility, **protected**, which makes a member visible to subclasses, but not to clients.
11.4.1.3 Constructors and Destructors

C++ allows the user to include constructor functions in class definitions, which are used to initialize the data members of newly created objects. A constructor may also allocate the heap-dynamic data that are referenced by the pointer members of the new object. Constructors are implicitly called when an object of the class type is created. A constructor has the same name as the class whose objects it initializes. Constructors can be overloaded, but of course each constructor of a class must have a unique parameter profile.

A C++ class can also include a function called a destructor, which is implicitly called when the lifetime of an instance of the class ends. As stated earlier, stack-dynamic class instances can contain pointer members that reference heap-dynamic data. The destructor function for such an instance can include a delete operator on the pointer members to deallocate the heap space they reference. Destructors are often used as a debugging aid, in which case they display or print the values of some or all of the object’s data members before those members are deallocated. The name of a destructor is the class’s name, preceded by a tilde (~).

Neither constructors nor destructors have return types, and neither use return statements. Both constructors and destructors can be explicitly called.

11.4.1.4 An Example

Our example of a C++ abstract data type is a stack:

```cpp
#include <iostream.h>
class Stack {
private:  //** These members are visible only to other
           //** members and friends (see Section 11.6.4)
    int *stackPtr;
    int maxLen;
    int topSub;
public:  //** These members are visible to clients
    Stack() {  //** A constructor
        stackPtr = new int [100];
        maxLen = 99;
        topSub = -1;
    }
    ~Stack() {delete [] stackPtr;}  //** A destructor
    void push(int number) {
        if (topSub == maxLen)
            cerr << "Error in push--stack is full\n";
        else stackPtr[++topSub] = number;
    }
    void pop() {
        if (empty())
            cerr << "Error in pop--stack is empty\n";
        else topSub--;
    }
    int top() {
```
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We discuss only a few aspects of this class definition, because it is not necessary to understand all of the details of the code. Objects of the Stack class are stack dynamic but include a pointer that references heap-dynamic data. The Stack class has three data members—stackPtr, maxLen, and topSub—all of which are private. stackPtr is used to reference the heap-dynamic data, which is the array that implements the stack. The class also has four public member functions—push, pop, top, and empty—as well as a constructor and a destructor. All of the member function definitions are included in this class, although they could have been externally defined. Because the bodies of the member functions are included, they are all implicitly inlined. The constructor uses the new operator to allocate an array of 100 int elements from the heap. It also initializes maxLen and topSub.

The following is an example program that uses the Stack abstract data type:

```c++
void main() {
    int topOne;
    Stack stk;  //** Create an instance of the Stack class
    stk.push(42);
    stk.push(17);
    topOne = stk.top();
    stk.pop();
    . . .
}
```

Following is a definition of the Stack class with only prototypes of the member functions. This code is stored in a header file with the .h file name extension. The definitions of the member functions follow the class definition. These use the scope resolution operator, ::, to indicate the class to which they belong. These definitions are stored in a code file with the file name extension .cpp.

```
// Stack.h - the header file for the Stack class
#include <iostream.h>
class Stack {
    private:  //** These members are visible only to other
              //** members and friends (see  Section 11.6.3)
        int *stackPtr;
        int maxLen;
        int topSub;
    public:   //** These members are visible to clients
        Stack();  //** A constructor
        ~Stack();  //** A destructor
        void push(int);
        void pop();
```
11.4 Language Examples

int top();

int empty();

// Stack.cpp - the implementation file for the Stack class
#include <iostream.h>
#include "Stack.h"
using std::cout;
Stack::Stack() {  //** A constructor
    stackPtr = new int [100];
    maxLen = 99;
    topSub = -1;
};  //** A destructor
void Stack::push(int number) {
    if (topSub == maxLen)
        cerr << "Error in push--stack is full\n";
    else
        stackPtr[++topSub] = number;
}

void Stack::pop() {
    if (topSub == -1)
        cerr << "Error in pop--stack is empty\n";
    else
        topSub--;
}

int top() {
    if (topSub == -1)
        cerr << "Error in top--stack is empty\n";
    else
        return (stackPtr[topSub]);
}

int Stack::empty() { return (topSub == -1); }

11.4.2 Abstract Data Types in Objective-C

As has been previously stated, Objective-C is similar to C++ in that its initial design was the C language with extensions to support object-oriented programming. One of the fundamental differences between the two is that Objective-C uses the syntax of Smalltalk for its method calls.

11.4.2.1 Encapsulation

The interface part of an Objective-C class is defined in a container called an interface with the following general syntax:

@interface class-name: parent-class {
    instance variable declarations
}

method prototypes
@end
The first and last lines, which begin with at signs (@), are directives.

The implementation of a class is packaged in a container naturally named `implementation`, which has the following syntax:

```plaintext
@implementation class-name
method definitions
@end
```

As in C++, in Objective-C classes are types.

Method prototypes have the following syntax:

```
(+ | -) (return-type) method-name [: (formal-parameters)] ;
```

When present, the plus sign indicates that the method is a class method; a minus sign indicates an instance method. The brackets around the formal parameters indicate that they are optional. Neither the parentheses nor the colon are present when there are no parameters. As in most other languages that support object-oriented programming, all instances of an Objective-C class share a single copy of its instance methods, but each instance has its own copy of the instance data.

The syntactic form of the formal parameter list is different from that of the more common languages, C, C++, Java, and C#. If there is one parameter, its type is specified in parentheses before the parameter’s name, as in the following method prototype:

```
-(void) meth1: (int) x;
```

This method’s name is `meth1:` (note the colon). A method with two parameters could appear as in the following example method prototype:

```
-(int) meth2: (int) x second: (float) y;
```

In this case, the method’s name is `meth2:second:`, although that is obviously a poorly chosen name. The last part of the name (second) could have been omitted, as in the following:

```
-(int) meth2: (int) x: (float) y;
```

In this case, the name of the method is `meth2::`

Method definitions are like method prototypes except that they have a brace-delimited sequence of statements in place of the semicolon.

The syntax of a call to a method with no parameters is as follows:

```
[object-name method-name] ;
```

If a method takes one parameter, a colon is attached to the method name and the parameter follows. There is no other punctuation between the method name and the parameter. For example, a call to a method named `add1` on the
object referenced by `myAdder` that takes one parameter, in this case the literal 7, would appear as follows:

```
[myAdder add1: 7];
```

If a method takes two parameters and has only one part to its name, a colon follows the first parameter and the second parameter follows that. No other punctuation is used between the two parameters. If there are more parameters, this pattern is repeated. For example, if `add1` takes three parameters and has no other parts to its name, it could be called with the following:

```
[myAdder add1: 7: 5: 3];
```

A method could have multiple parameters and multiple parts to its name, as in the previous example:

```
-(int) meth2: (int) x second: (float) y;
```

An example call to this method follows:

```
[myObject meth2: 7 second: 3.2];
```

Constructors in Objective-C are called `initializers`; they only provide initial values. They can be given any name, and as a result they must be explicitly called. Constructors return a reference to the new object, so their type is always a pointer to the class name. They use a `return` statement that returns `self`, a reference to the current object.

An object is created in Objective-C by calling `alloc`. Typically, after calling `alloc`, the constructor of the class is explicitly called. These two calls can be and usually are cascaded, as in the following statement, which creates an object of `Adder` class with `alloc` and then calls its constructor, `init`, on the new object, and puts the address of the new object in `myAdder`:

```
Adder *myAdder = [[Adder alloc] init];
```

All class instances are heap dynamic and are referenced through reference variables.

C programs nearly always import a header file for input and output functions, `stdio.h`. In Objective-C, a header file is usually imported that has the prototypes of a variety of often required functions, including those for input and output, as well as some needed data. This is done with the following:

```
#import <Foundation/Foundation.h>
```

Nearly all of the code in the `main` function of a program usually is placed in a compound statement that follows `@autoreleasepool`. Any objects that are created in the compound statement are automatically released at the end of the compound statement.
The `@autoreleasepool` compound statement in a `main` function is followed by a `return 0` statement, indicating that execution successfully reached the end of `main`'s code.

### 11.4.2.2 Information Hiding

Objective-C uses the directives, `@private` and `@public`, to specify the access levels of the instance variables in a class definition. These are used as the reserved words `public` and `private` as used in C++. The difference is that the default in Objective-C is protected, whereas it is private in C++. Unlike most programming languages that support object-oriented programming, in Objective-C there is no way to restrict access to a method.

In Objective-C, the convention is that the name of a getter method for an instance variable is the variable's name. The name of the setter method is the word `set` with the capitalized variable's name attached. So, for a variable named `sum`, the getter method would be named `sum` and the setter method would be named `setSum`. Assuming that `sum` is an `int` variable, these methods could be defined as follows:

```objective-c
// The getter for sum
-(int) sum {
    return sum;
}

// The setter for sum
-(void) setSum: (int) s {
    sum = s;
}
```

If the getter and setter method for a particular variable does not impose any constraints on their actions, they can be automatically generated by the Objective-C compiler. This is done by listing the instance variables for which getters and setters are to be generated on the property directive in the interface section, as in the following:

```objective-c
@property int sum;
```

In the implementation section, the variables are listed in a `synthesize` directive, as in the following:

```objective-c
@synthesize sum;
```

Variables for which getters and setters are generated by the compiler are often called `properties` and the accessor methods are said to be `synthesized`. The getters and setters of instance variables can be used in two ways, either in method calls or in dot notation, as if they were publically accessible. For example, if we have defined a getter and a setter for the variable `sum`, they could be used as in the following:
[myObject setSum: 100];
newSum = [myObject sum];

or as if they were publically accessible, as in the following:

myObject.sum = 100;
newSum = myObject.sum;

11.4.2.3 An Example

Following are the definitions of the interface and implementation of the stack class in Objective-C:

// stack.m - interface and implementation of a simple stack
#import <Foundation/Foundation.h>

// Interface section
@interface Stack: NSObject {
    int stackArray [100];
    int stackPtr;
    int maxLen;
    int topSub;
}
-(void) push: (int) number;
-(void) pop;
-(int) top;
-(int) empty;
@end

// Implementation section

@implementation Stack
-(Stack *) initWith {
    maxLen = 100;
    topSub = -1;
    stackPtr = stackArray;
    return self;
}

-(void) push: (int) number {
    if (topSub == maxLen)
        NSLog(@"Error in push--stack is full");
    else
        stackPtr[++] = number;
}

-(void) pop {
    if (topSub == -1)
NSLog(@"Error in pop--stack is empty");
else
    topSub--;
}

-(int) top {
    if (topSub >= 0)
        return stackPtr[topSub];
    else
        NSLog(@"Error in top--stack is empty");
}

-(int) empty {
    return topSub == -1);
}

int main (int argc, char *argv[]) { 
    int temp;
    @autoreleasepool{
    Stack *myStack = [[Stack alloc]initWith];
    [myStack push: 5];
    [myStack push: 3];
    temp = [myStack top];
    NSLog(@"Top element is:%i", temp);
    [myStack pop];
    temp = [myStack top];
    NSLog(@"Top element is:%i", temp);
    temp = [myStack top];
    [myStack pop];
    }
    return 0;
}
@end

The output of this program is as follows:

Top element is: 3
Top element is: 5
Error in top--stack is empty
Error in pop--stack is empty

Screen output from an Objective-C program is created with a call to a method with the odd-looking name, NSLog, which takes a literal string as its parameter. Literal strings are created with an at sign (@) followed by a quoted string. If an output string includes the values of variables, the names of the variables are included as parameters in the call to NSLog. The positions in the literal string for the values are marked with format codes, for example %i for an integer and %f for a floating-point value in scientific notation, as is similar to C's printf function.
11.4.2.4 Evaluation

The support in Objective-C for abstract data types is adequate. Some find it odd that it uses syntactic forms from two very different languages, Smalltalk (for its method calls) and C (for nearly everything else). Also, its use of directives in place of language constructs to indicate class interfaces and implementation sections also differs from most other programming languages. One minor deficiency is the lack of a way to restrict access to methods. So, even methods meant only to be used inside a class are accessible to clients. Another minor deficiency is that constructors must be explicitly called, thereby requiring programmers to remember to call them, and also leading to further clutter in the client program.

11.4.3 Abstract Data Types in Java

Java support for abstract data types is similar to that of C++. There are, however, a few important differences. All objects are allocated from the heap and accessed through reference variables. Methods in Java must be defined completely in a class. A method body must appear with its corresponding method header. Therefore, a Java abstract data type is both declared and defined in a single syntactic unit. A Java compiler can inline any method that is not overridden. Definitions are hidden from clients by declaring them to be private.

One important advantage of Java’s classes over the classes of C++ and Objective-C is that it uses implicit garbage collection of all objects. This allows the programmer to ignore the issue of deallocation of objects and the clutter of deallocation code in the implementations of abstract data types.

Rather than having private and public clauses in its class definitions, in Java access modifiers can be attached to method and variable definitions. If an instance variable or method does not have an access modifier, it has package access, which is discussed in Section 11.7.2.

11.4.3.1 An Example

The following is a Java class definition for our stack example:

```java
class StackClass {
    private int[] stackRef;
    private int maxLen,
        topIndex;
    public StackClass() { // A constructor
        stackRef = new int[100];
        maxLen = 99;
    }
}
```

---

2. Java interfaces are an exception to this—an interface has method headers but cannot include their bodies.
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topIndex = -1;
}
public void push(int number) {
    if (topIndex == maxLen)
        System.out.println("Error in push–stack is full");
    else stackRef[++topIndex] = number;
}
public void pop() {
    if (empty())
        System.out.println("Error in pop–stack is empty");
    else --topIndex;
}
public int top() {
    if (empty()) {
        System.out.println("Error in top–stack is empty");
        return 9999;
    }
    else
        return (stackRef[topIndex]);
}
public boolean empty() { return (topIndex == -1); }
}

An example class that uses StackClass follows:

public class TstStack {
    public static void main(String[] args) {
        StackClass myStack = new StackClass();
        myStack.push(42);
        myStack.push(29);
        System.out.println("29 is: " + myStack.top());
        myStack.pop();
        System.out.println("42 is: " + myStack.top());
        myStack.pop();
        myStack.pop();  // Produces an error message
    }
}

One obvious difference is the lack of a destructor in the Java version, obviated by Java’s implicit garbage collection.3

11.4.3.2 Evaluation

Although different in some primarily cosmetic ways, Java’s support for abstract data types is similar to that of C++. Java clearly provides for what is necessary to design abstract data types.

3. In Java, the finalize method serves as a kind of destructor.
11.4 Language Examples

11.4.4 Abstract Data Types in C#

Recall that C# is based on both C++ and Java and that it also includes some new constructs. Like Java, all C# class instances are heap dynamic. Default constructors, which provide initial values for instance data, are predefined for all classes. These constructors provide typical initial values, such as 0 for int types and false for boolean types. A user can furnish one or more constructors for any class he or she defines. Such constructors can assign initial values to some or all of the instance data of the class. Any instance variable that is not initialized in a user-defined constructor is assigned a value by the default constructor.

Although C# allows destructors to be defined, because it uses garbage collection for most of its heap objects, destructors are rarely used.

11.4.4.1 Encapsulation

As mentioned in Section 11.4.2, C++ includes both classes and structs, which are nearly identical constructs. The only difference is that the default access modifier for class is private, whereas for structs it is public. C# also has structs, but they are very different from those of C++. In C#, structs are, in a sense, lightweight classes. They can have constructors, properties, methods, and data fields and can implement interfaces but do not support inheritance. One other important difference between structs and classes in C# is that structs are value types, as opposed to reference types. They are allocated on the run-time stack, rather than the heap. If they are passed as parameters, like other value types, by default they are passed by value. All C# value types, including all of its primitive types, are actually structs. Structs can be created by declaring them, like other predefined value types, such as int or float. They can also be created with the new operator, which calls a constructor to initialize them.

Structs are used in C# primarily to implement relatively small simple types that need never be base types for inheritance. They are also used when it is convenient for the objects of the type to be stack as opposed to heap allocated.

11.4.4.2 Information Hiding

C# uses the private and protected access modifiers exactly as they are used in Java.

C# provides properties, which it inherited from Delphi, as a way of implementing getters and setters without requiring explicit method calls by the client. As with Objective-C, properties provide implicit access to specific private instance data. For example, consider the following simple class and client code:

```csharp
generics

class Weather
{
    public int DegreeDays
    { //** DegreeDays is a property
        get
        {
            return degreeDays;
        }
    }
}
set {
    if (value < 0 || value > 30)
        Console.WriteLine(
            "Value is out of range: {0}", value);
    else
        degreeDays = value;
}

private int degreeDays;
...
Weather w = new Weather();
int degreeDaysToday, oldDegreeDays;
...
w.DegreeDays = degreeDaysToday;
...
oldDegreeDays = w.DegreeDays;

In the class Weather, the property DegreeDays is defined. This property provides a getter method and a setter method for access to the private data member, degreeDays. In the client code following the class definition, degreeDays is treated as if it were a public-member variable, although access to it is available through the property only. Notice the use of the implicit variable value in the setter method. This is the mechanism by which the new value of the property is referenced.

The stack example is not shown here in C#. The only difference between the Java version in Section 11.4.4.1 and the C# version is the output method calls and the use of bool instead of boolean for the return type of the empty method.

11.4.5 Abstract Data Types in Ruby

Ruby provides support for abstract data types through its classes. In terms of capabilities, Ruby classes are similar to those in C++ and Java.

11.4.5.1 Encapsulation

In Ruby, a class is defined in a compound statement opened with the class reserved word and closed with end. The names of instance variables have a special syntactic form, they must begin with at signs (@). Instance methods have the same syntax as functions in Ruby: They begin with the def reserved word and end with end. Class methods are distinguished from instance methods by having the class name appended to the beginning of their names with a period separator. For example, in a class named Stack, a class method's name would begin with Stack. Constructors in Ruby are named initialize. Because the constructor cannot be overloaded, there only can be one per class.
Classes in Ruby are dynamic in the sense that members can be added at any time. This is done by simply including additional class definitions that specify the new members. Moreover, even predefined classes of the language, such as `String`, can be extended. For example, consider the following class definition:

```ruby
class myClass
  def meth1
    . . .
  end
end
```

This class could be extended by adding a second method, `meth2`, with a second class definition:

```ruby
class myClass
  def meth2
    . . .
  end
end
```

Methods can also be removed from a class. This is done by providing another class definition in which the method to be removed is sent to the method `remove_method` as a parameter. The dynamic classes of Ruby is another example of a language designer trading readability (and as a consequence, reliability) for flexibility. Allowing dynamic changes to classes clearly adds flexibility to the language, while harming readability. To determine the behavior of a class at a particular point in a program, one must find all of its definitions in the program and consider all of them.

### 11.4.5.2 Information Hiding

Access control for methods in Ruby is dynamic, so access violations are detected only during execution. The default method access is public, but it can also be protected or private. There are two ways to specify the access control, both of which use functions with the same names as the access levels, `private` and `public`. One way is to call the appropriate function without parameters. This resets the default access for subsequently defined methods in the class. For example,
The alternative is to call the access control functions with the names of the specific methods as parameters. For example, the following is semantically equivalent to the previous class definition:

```ruby
class MyClass
  def meth1
    ...
  end
  ...
  def meth7
    ...
  end
  private :meth7, ...
end
```

In Ruby, all data members of a class are private, and that cannot be changed. So, data members can be accessed only by the methods of the class, some of which may be accessor methods. In Ruby, instance data that are accessible through accessor methods are called attributes.

For an instance variable named `@sum`, the getter and setter methods would be as follows:

```ruby
def sum
  @sum
end
def sum=(new_sum)
  @sum = new_sum
end
```

Notice that getters are given the name of the instance variable minus the `@`. The names of setter methods are the same as those of the corresponding getters, except they have an equal sign (`=`) attached.

Getters and setters can be implicitly generated by the Ruby system by including calls to `attr_reader` and `attr_writer`, respectively, in the class definition. The parameters to these are the symbols of the attribute’s names, as is illustrated in the following:

```ruby
attr_reader :sum, :total
attr_writer :sum
```

### 11.4.5.3 An Example

Following is the stack example written in Ruby:

```ruby
# Stack.rb - defines and tests a stack of maximum length 100, implemented in an array
```
class StackClass

# Constructor
  def initialize
    @stackRef = Array.new(100)
    @maxLen = 100
    @topIndex = -1
  end

# push method
  def push(number)
    if @topIndex == @maxLen
      puts "Error in push - stack is full"
    else
      @topIndex = @topIndex + 1
      @stackRef[@topIndex] = number
    end
  end

# pop method
  def pop
    if empty
      puts "Error in pop - stack is empty"
    else
      @topIndex = @topIndex - 1
    end
  end

# top method
  def top
    if empty
      puts "Error in top - stack is empty"
    else
      @stackRef[@topIndex]
    end
  end

# empty method
  def empty
    @topIndex == -1
  end
end  # of Stack class

# Test code for StackClass
myStack = StackClass.new
myStack.push(42)
myStack.push(29)
puts "Top element is (should be 29): #{myStack.top}"
myStack.pop
puts "Top element is (should be 42): #{myStack.top}"
myStack.pop
Recall that the notation \{variable\} converts the value of the variable to a string, which is then inserted into the string in which it appears. This class defines a stack structure that can store objects of any type.

### 11.4.5.4 Evaluation

Recall that in Ruby, everything is an object and arrays are actually arrays of references to objects. That clearly makes this stack more flexible than the similar examples in C++ and Java. Furthermore, simply by passing the desired maximum length to the constructor, objects of this class could have any given maximum length. Of course, because arrays in Ruby have dynamic length, the class could be modified to implement stack objects that are not restricted to any length, except that imposed by the machine's memory capacity. Because the names of class and instance variables have different forms, Ruby has a slight readability advantage over the other languages discussed in this section.

### 11.5 Parameterized Abstract Data Types

It is often convenient to be able to parameterize abstract data types. For example, we should be able to design a stack abstract data type that can store any scalar type elements rather than be required to write a separate stack abstraction for every different scalar type. Note that this is only an issue for static typed languages. In a dynamic typed language like Ruby, any stack implicitly can store any type elements. In fact, different elements of the stack could be of different types. In the following three subsections, the capabilities of C++, Java 5.0, and C# 2005 to construct parameterized abstract data types are discussed.

#### 11.5.1 C++

To make the example C++ stack class of Section 11.4.1 generic in the stack size, only the constructor function needs to be changed, as in the following:

```cpp
Stack(int size) {
    stackPtr = new int [size];
    maxLen = size - 1;
    topSub = -1;
}
```

The declaration for a stack object now may appear as follows:

```cpp
Stack stk(150);
```
The class definition for Stack can include both constructors, so users can use the default-size stack or specify some other size.

The element type of the stack can be made generic by making the class a templated class. Then, the element type can be a template parameter. The definition of the templated class for a stack type is as follows:

```cpp
#include <iostream.h>
template <typename Type>  // Type is the template parameter
class Stack {
    private:
        Type *stackPtr;
        int maxLen;
        int topSub;
    public:
        // A constructor for 100 element stacks
        Stack() {
            stackPtr = new Type [100];
            maxLen = 99;
            topSub = -1;
        }
        // A constructor for a given number of elements
        Stack(int size) {
            stackPtr = new Type [size];
            maxLen = size - 1;
            topSub = -1;
        }
        ~Stack() {delete stackPtr;};  // A destructor
        void push(Type number) {
            if (topSub == maxLen)
                cout << "Error in push–stack is full\n";
            else stackPtr[++ topSub] = number;
        }
        void pop() {
            if (empty())
                cout << "Error in pop–stack is empty\n";
            else topSub --;
        }
        Type top() {
            if (empty())
                cerr << "Error in top--stack is empty\n";
            else return (stackPtr[topSub]);
        }
        int empty() {return (topSub == -1);} 
    }

C++ templated classes are instantiated to become typed classes at compile time. For example, an instance of the templated Stack class, as well as an instance of the typed class, can be created with the following declaration:

```cpp
Stack<int> myIntStack;
```
However, if an instance of the templated `Stack` class has already been created for the `int` type, the typed class need not be created.

### 11.5.2 Java 5.0

Java 5.0 supports a form of parameterized abstract data types in which the generic parameters must be classes. Recall that these were briefly discussed in Chapter 9.

The most common generic types are collection types, such as `LinkedList` and `ArrayList`, which were in the Java class library before support for generics was added. The original collection types stored `Object` class instances, so they could store any objects (but not primitive types). Therefore, the collection types have always been able to store multiple types (as long as they are classes). There were three issues with this: First, every time an object was removed from the collection, it had to be cast to the appropriate type. Second, there was no error checking when elements were added to the collection. This meant that once the collection was created, objects of any class could be added to the collection, even if the collection was meant to store only `Integer` objects. Third, the collection types could not store primitive types. So, to store `int` values in an `ArrayList`, the value first had to be put in an `Integer` class instance. For example, consider the following code:

```java
/* Create an ArrayList object
ArrayList myArray = new ArrayList();
/* Create an element
myArray.add(0, new Integer(47));
/* Get first object
Integer myInt = (Integer)myArray.get(0);
```

In Java 5.0, the collection classes, the most commonly used of which is `ArrayList`, became a generic class. Such classes are instantiated by calling `new` on the class constructor and passing it the generic parameter in pointed brackets. For example, the `ArrayList` class can be instantiated to store `Integer` objects with the following statement:

```java
ArrayList <Integer> myArray = new ArrayList <Integer>();
```

This new class overcomes two of the problems with pre-Java 5.0 collections. Only `Integer` objects can be put into the `myArray` collection. Furthermore, there is no need to cast an object being removed from the collection.

Java 5.0 also includes generic collections for linked-lists, queues, and sets. Users also can define generic classes in Java 5.0. For example, we could have the following:

```java
public class MyClass<T> {  
  . . .
}
```
This class could be instantiated with the following:

```
MyClass<String> myString;
```

There are some drawbacks to these user-defined generic classes. For one thing, they cannot store primitives. Second, the elements cannot be indexed. Elements must be added to user-defined generic collections with the `add` method. Next, we implement the generic stack example using an `ArrayList`. Note that the last element of an `ArrayList` is found using the `size` method, which returns the number of elements in the structure. Elements are deleted from the structure with the `remove` method. Following is the generic class:

```
import java.util.*;
public class Stack2<T> {

  private ArrayList<T> stackRef;
  private int maxLen;

  public Stack2() { // A constructor
    stackRef = new ArrayList<T> ();
    maxLen = 99;
  }

  public void push(T newValue) {
    if (stackRef.size() == maxLen)
      System.out.println("Error in push–stack is full");
    else
      stackRef.add(newValue);
  }

  public void pop() {
    if (empty())
      System.out.println("Error in pop–stack is empty");
    else
      stackRef.remove(stackRef.size() - 1);
  }

  public T top() {
    if (empty()) {
      System.out.println("Error in top–stack is empty");
      return null;
    } else
      return (stackRef.get(stackRef.size() - 1));
  }

  public boolean empty() { return (stackRef.isEmpty());}

This class could be instantiated for the `String` type with the following:

```
Stack2<String> myStack = new Stack2<String>();
```

Recall from Chapter 9, that Java 5.0 supports wildcard classes. For example, `Collection<?>` is a wildcard class for all collection classes. This allows a method to be written that can accept any collection type as a parameter. Because a collection can itself be generic, the `Collection<?>` class is in a sense a generic of a generic class.
Some care must be taken with objects of the wildcard type. For example, because the components of a particular object of this type have a type, other type objects cannot be added to the collection. For example, consider

```
Collection<?> c = new ArrayList<String>();
```

It would be illegal to use the `add` method to put something into this collection unless its type were `String`.

A generic class can easily be defined in Java 5.0 that will work only for a restricted set of types. For example, a class can declare a variable of the generic type and call a method such as `compareTo` through that variable. If the class is instantiated for a type that does not include a `compareTo` method, the class cannot be used. To prevent a generic class from being instantiated for a type that does not support `compareTo`, it could be defined with the following generic parameter:

```
<T extends Comparable>
```

`Comparable` is the interface in which `compareTo` is declared. If this generic type is used on a class definition, the class cannot be instantiated for any type that does not implement `Comparable`. The choice of the reserved word `extends` seems odd here, but its use is related to the concept of a subtype. Apparently, the designers of Java did not want to add another more connotative reserved word to the language.

### 11.5.3 C# 2005

As was the case with Java, the first version of C# defined collection classes that stored objects of any class. These were `ArrayList`, `Stack`, and `Queue`. These classes had the same problems as the collection classes of pre-Java 5.0.

Generic classes were added to C# in its 2005 version. The five predefined generic collections are `Array`, `List`, `Stack`, `Queue`, and `Dictionary` (the `Dictionary` class implements hashes). Exactly as in Java 5.0, these classes eliminate the problems of allowing mixed types in collections and requiring casts when objects are removed from the collections.

As with Java 5.0, users can define generic classes in C# 2005. One capability of the user-defined C# generic collections is that any of them can be defined to allow its elements to be indexed (accessed through subscripting). Although the indexes are usually integers, an alternative is to use strings as indexes.

One capability that Java 5.0 provides that C# 2005 does not is wildcard classes.

### 11.6 Encapsulation Constructs

The first five sections of this chapter discussed abstract data types, which are minimal encapsulations. This section describes the multiple-type encapsulations that are needed for larger programs.
11.6.1 Introduction

When the size of a program reaches beyond a few thousand lines, two practical problems become evident. From the programmer’s point of view, having such a program appear as a single collection of subprograms or abstract data type definitions does not impose an adequate level of organization on the program to keep it intellectually manageable. The second practical problem for larger programs is recompilation. For relatively small programs, recompiling the whole program after each modification is not costly. But for large programs, the cost of recompilation is significant. So, there is an obvious need to find ways to avoid recompilation of the parts of a program that are not affected by a change. The obvious solution to both of these problems is to organize programs into collections of logically related code and data, each of which can be compiled without recompilation of the rest of the program. An encapsulation is such a collection.

Encapsulations are often placed in libraries and made available for reuse in programs other than those for which they were written. People have been writing programs with more than a few thousand lines for at least the last 50 years, so techniques for providing encapsulations have been evolving for some time.

In languages that allow nested subprograms, programs can be organized by nesting subprogram definitions inside the logically larger subprograms that use them. This can be done in Python and Ruby. As discussed in Chapter 5, however, this method of organizing programs, which uses static scoping, is far from ideal. Therefore, even in languages that allow nested subprograms, they are not used as a primary organizing encapsulation construct.

11.6.2 Encapsulation in C

C does not provide complete support for abstract data types, although both abstract data types and multiple-type encapsulations can be simulated.

In C, a collection of related functions and data definitions can be placed in a file, which can be independently compiled. Such a file, which acts as a library, has an implementation of its entities. The interface to such a file, including data, type, and function declarations, is placed in a separate file called a header file. Type representations can be hidden by declaring them in the header file as pointers to struct types. The complete definitions of such struct types need only appear in the implementation file.

The header file, in source form, and the compiled version of the implementation file are furnished to clients. When such a library is used, the header file is included in the client code, using an #include preprocessor specification, so that references to functions and data in the client code can be type checked. The #include specification also documents the fact that the client program depends on the library implementation file. This approach effectively separates the specification and implementation of an encapsulation.

Although these encapsulations work, they create some insecurities. For example, a user could simply cut and paste the definitions from the header file.
into the client program, rather than using \texttt{#include}. This would work, because \texttt{#include} simply copies the contents of its operand file into the file in which the \texttt{#include} appears. However, there are two problems with this approach. First, the documentation of the dependence of the client program on the library (and its header file) is lost. Second, suppose a user copies the header file into his or her program. Then the author of the library changes both the header and the implementation files. Following this, the user uses the new implementation file with the old header. For example, a variable $x$ could have been defined to be \texttt{int} type in the old header file, which the client code still uses, although the implementation code has been recompiled with the new header file, which defines $x$ to be \texttt{float}. So, the implementation code was compiled with $x$ as an \texttt{int} but the client code was compiled with $x$ as a \texttt{float}. The linker does not detect this error.

Thus, it is the user’s responsibility to ensure that both the header and implementation files are up-to-date. This is often done with a \texttt{make} utility.

One other drawback of this approach is the inherent problems of pointers and the potential confusion with assignment and comparisons of pointers.

11.6.3 Encapsulation in C++

C++ provides two different kinds of encapsulation: header and implementation files can be defined as in C, or class headers and definitions can be defined. Because of the complex interplay of C++ templates and separate compilation, the header files of C++ template libraries often include complete definitions of resources, rather than just data declarations and subprogram protocols; this is due in part to the use of the C linker for C++ programs.

When nontemplated classes are used for encapsulations, the class header file has only the prototypes of the member functions, with the function definitions provided outside the class in a code file, as in the last example in Section 11.4.1.4. This clearly separates interface from implementation.

One language design problem that results from having classes but no generalized encapsulation construct is that sometimes when operations are defined that use two different classes of objects, the operation does not naturally belong in either class. For example, suppose we have an abstract data type for matrices and one for vectors and need a multiplication operation between a vector and a matrix. The multiplication code must have access to the data members of both the vector and the matrix classes, but neither of those classes is the natural home for the code. Furthermore, regardless of which is chosen, access to the members of the other is a problem. In C++, these kinds of situations can be handled by allowing nonmember functions to be “friends” of a class. Friend functions have access to the private entities of the class where they are declared to be friends. For the matrix/vector multiplication operation, one C++ solution is to define the operation outside both the matrix and the vector classes but define it to be a friend of both. The following skeletal code illustrates this scenario:
class Matrix;  //** A class declaration
class Vector {
    friend Vector multiply(const Matrix&, const Vector&);
    . . .
};
class Matrix {  //** The class definition
    friend Vector multiply(const Matrix&, const Vector&);
    . . .
};
/** The function that uses both Matrix and Vector objects
Vector multiply(const Matrix& m1, const Vector& v1) {
    . . .
}

In addition to functions, whole classes can be defined to be friends of a class; then all the private members of the class are visible to all of the members of the friend class.

11.6.4 C# Assemblies

C# includes a larger encapsulation construct than a class. The construct is the one used by all of the .NET programming languages: the assembly. Assemblies are built by .NET compilers. A .NET application consists of one or more assemblies. An assembly is a file\(^4\) that appears to application programs to be a single dynamic link library (.dll)\(^5\) or an executable (.exe). An assembly defines a module, which can be separately developed. An assembly includes several different components. One of the primary components of an assembly is its programming code, which is in an intermediate language, having been compiled from its source language. In .NET, the intermediate language is named Common Intermediate Language (CIL). It is used by all .NET languages. Because its code is in CIL, an assembly can be used on any architecture, device, or operating system. When executed, the CIL is just-in-time compiled to native code for the architecture on which it is resident.

In addition to the CIL code, a .NET assembly includes metadata that describes every class it defines, as well as all external classes it uses. An assembly also includes a list of all assemblies referenced in the assembly and an assembly version number.

\(^4\) An assembly can consist of any number of files.

\(^5\) A dynamic link library (DLL) is a collection of classes and methods that are individually linked to an executing program when needed during execution. Therefore, although a program has access to all of the resources in a particular DLL, only the parts that are actually used are ever loaded and linked to the program. DLLs have been part of the Windows programming environment since Windows first appeared. However, the DLLs of .NET are quite different from those of previous Windows systems.
In the .NET world, the assembly is the basic unit of deployment of software. Assemblies can be private, in which case they are available to just one application, or public, which means any application can use them.

As mentioned previously, C# has an access modifier, internal. An internal member of a class is visible to all classes in the assembly in which it appears.

Java has a file structure that is similar to an assembly called a Java Archive (JAR). It is also used for deployment of Java software systems. JARs are built with the Java utility jar, rather than a compiler.

### 11.7 Naming Encapsulations

We have considered encapsulations to be syntactic containers for logically related software resources—in particular, abstract data types. The purpose of these encapsulations is to provide a way to organize programs into logical units for compilation. This allows parts of programs to be recompiled after isolated changes. There is another kind of encapsulation that is necessary for constructing large programs: a naming encapsulation.

A large program is usually written by many developers, working somewhat independently, perhaps even in different geographic locations. This requires the logical units of the program to be independent, while still able to work together. It also creates a naming problem: How can independently working developers create names for their variables, methods, and classes without accidentally using names already in use by some other programmer developing a different part of the same software system?

Libraries are the origin of the same kind of naming problems. Over the past two decades, large software systems have become progressively more dependent on libraries of supporting software. Nearly all software written in contemporary programming languages requires the use of large and complex standard libraries, in addition to application-specific libraries. This widespread use of multiple libraries has necessitated new mechanisms for managing names. For example, when a developer adds new names to an existing library or creates a new library, he or she must not use a new name that conflicts with a name already defined in a client’s application program or in some other library. Without some language processor assistance, this is virtually impossible, because there is no way for the library author to know what names a client’s program uses or what names are defined by the other libraries the client program might use.

Naming encapsulations define name scopes that assist in avoiding these name conflicts. Each library can create its own naming encapsulation to prevent its names from conflicting with the names defined in other libraries or in client code. Each logical part of a software system can create a naming encapsulation with the same purpose.

Naming encapsulations are logical encapsulations, in the sense that they need not be physically contiguous. Several different collections of code can be placed in the same namespace, even though they are stored in different places. In the following sections, we briefly describe the uses of naming encapsulations in C++, Java, and Ruby.
11.7.1 C++ Namespaces

C++ includes a specification, `namespace`, that helps programs manage the problem of global namespaces. One can place each library in its own namespace and qualify the names in the program with the name of the namespace when the names are used outside that namespace. For example, suppose there is an abstract data type header file that implements stacks. If there is concern that some other library file may define a name that is used in the stack abstract data type, the file that defines the stack could be placed in its own namespace. This is done by placing all of the declarations for the stack in a namespace block, as in the following:

```cpp
namespace myStackSpace {
    // Stack declarations
}
```

The implementation file for the stack abstract data type could reference the names declared in the header file with the scope resolution operator, `::`, as in

```cpp
myStackSpace::topSub
```

The implementation file could also appear in a namespace block specification identical to the one used on the header file, which would make all of the names declared in the header file directly visible. This is definitely simpler, but slightly less readable, because it is less obvious where a specific name in the implementation file is declared.

Client code can gain access to the names in the namespace of the header file of a library in three different ways. One way is to qualify the names from the library with the name of the namespace. For example, a reference to the variable `topSub` could appear as follows:

```cpp
myStackSpace::topSub
```

This is exactly the way the implementation code could reference it if the implementation file was not in the same namespace.

The other two approaches use the `using` directive. This directive can be used to qualify individual names from a namespace, as with

```cpp
using myStackSpace::topSub;
```

which makes `topSub` visible, but not any other names from the `myStackSpace` namespace.

The `using` directive also can be used to qualify all of the names from a namespace, as in the following:

```cpp
using namespace myStackSpace;
```

Code that includes this directive can directly access the names defined in the namespace, as in

```cpp
p = topSub;
```
Be aware that namespaces are a complicated feature of C++, and we have introduced only the simplest part of the story here.

C# includes namespaces that are much like those of C++.

11.7.2 Java Packages

Java includes a naming encapsulation construct: the package. Packages can contain more than one type\(^6\) definition, and the types in a package are partial friends of one another. Partial here means that the entities defined in a type in a package that either are public or protected (see Chapter 12) or have no access specifier are visible to all other types in the package.

Entities without access modifiers are said to have package scope, because they are visible throughout the package. Java therefore has less need for explicit friend declarations and does not include the friend functions or friend classes of C++.

The resources defined in a file are specified to be in a particular package with a package declaration, as in

```java
class package stkpkg;
```

The package declaration must appear as the first line of the file. The resources of every file that does not include a package declaration are implicitly placed in the same unnamed package.

The clients of a package can reference the types defined in the package using fully qualified names. For example, if the package `stkpkg` has a class named `myStack`, that class can be referenced in a client of `stkpkg` as `stkpkg.myStack`. Likewise, a variable in the `myStack` object named `topSub` could be referenced as `stkpkg.myStack.topSub`. Because this approach can quickly become cumbersome when packages are nested, Java provides the `import` declaration, which allows shorter references to type names defined in a package. For example, suppose the client includes the following:

```java
import stkpkg.myStack;
```

Now, the class `myStack` can be referenced by just its name. To be able to access all of the type names in the package, an asterisk can be used on the import statement in place of the type name. For example, if we wanted to import all of the types in `stkpkg`, we could use the following:

```java
import stkpkg.*;
```

Note that Java’s `import` is only an abbreviation mechanism. No otherwise hidden external resources are made available with `import`. In fact, in Java nothing is implicitly hidden if it can be found by the compiler or class loader (using the package name and the `CLASSPATH` environment variable).

---

6. By type here we mean either a class or an interface.
Java’s `import` documents the dependencies of the package in which it appears on the packages named in the `import`. These dependencies are less obvious when `import` is not used.

### 11.7.3 Ruby Modules

Ruby classes serve as namespace encapsulations, as do the classes of other languages that support object-oriented programming. Ruby has an additional naming encapsulation, called a module. Modules typically define collections of methods and constants. So, modules are convenient for encapsulating libraries of related methods and constants, whose names are in a separate namespace so there are no name conflicts with other names in a program that uses the module. Modules are unlike classes in that they cannot be instantiated or subclassed and do not define variables. Methods that are defined in a module include the module’s name in their names. For example, consider the following skeletal module definition:

```ruby
module MyStuff
  PI = 3.14159265
  def MyStuff.mymethod1(p1)
    ...
  end
  def MyStuff.mymethod2(p2)
    ...
  end
end
```

Assuming the `MyStuff` module is stored in its own file, a program that wants to use the constant and methods of `MyStuff` must first gain access to the module. This is done with the `require` method, which takes the file name in the form of a string literal as a parameter. Then, the constants and methods of the module can be accessed through the module’s name. Consider the following code that uses our example module, `MyStuff`, which is stored in the file named `myStuffMod`

```ruby
require 'myStuffMod'
...
MyStuff.mymethod1(x)
...
```

Modules are further discussed in Chapter 12.

---

**Summary**

The concept of abstract data types, and their use in program design, was a milestone in the development of programming as an engineering discipline. Although the concept is relatively simple, its use did not become convenient and safe until languages were designed to support it.
The two primary features of abstract data types are the packaging of data objects with their associated operations and information hiding. A language may support abstract data types directly or simulate them with more general encapsulations.

C++ data abstraction is provided by classes. Classes are types, and instances can be either stack or heap dynamic. A member function (method) can have its complete definition appear in the class or have only the protocol given in the class and the definition placed in another file, which can be separately compiled. C++ classes can have two clauses, each prefixed with an access modifier: private or public. Both constructors and destructors can be given in class definitions. Heap-allocated objects must be explicitly deallocated with `delete`.

As with C++, Objective-C data abstractions are classes. Classes are types and all are heap dynamic. Methods declarations must appear in interface sections of classes and method definitions must appear in implementation sections. Constructors are called `initializers`; they must be explicitly called. Instance variables can be private or public. Access to methods cannot be restricted. Method calls use syntax that is similar to that used by Smalltalk. Objective-C supports properties and access methods for properties can be furnished by the compiler.

Java data abstractions are similar to those of C++, except all Java objects are allocated from the heap and are accessed through reference variables. Also, all objects are garbage collected. Rather than having access modifiers attached to clauses, in Java the modifiers appear on individual declarations (or definitions).

C# supports abstract data types with both classes and structs. Its structs are value types and do not support inheritance. C# classes are similar to those of Java.

Ruby supports abstract data types with its classes. Ruby’s classes differ from those of most other languages in that they are dynamic—members can be added, deleted, or changed during execution.

C++, Java 5.0, and C# 2005 allow their abstract data types to be parameterized—Ada through its generic packages, C++ through its templated classes, and Java 5.0 and C# through their collection classes and interfaces and user-defined generic classes.

To support the construction of large programs, some contemporary languages include multiple-type encapsulation constructs, which can contain a collection of logically related types. An encapsulation may also provide access control to its entities. Encapsulations provide the programmer with a method of organizing programs that also facilitates recompilation.

C++, C#, Java, and Ruby provide naming encapsulations. For Ada and Java, they are named packages; for C++ and C#, they are namespaces; for Ruby, they are modules. Partially because of the availability of packages, Java does not have friend functions or friend classes.

**Review Questions**

1. What is an abstraction?
2. Define object.
3. What is the standard for floating-point number representation?
4. Name two accessor methods that allow indirect access to hidden data.
5. What are the languages that support abstract data types?
6. What is the use of a C++ destructor?
7. What are called “initializers” in Objective-C?
8. How does Objective-C use information-hiding directives?
9. What is the advantage of the classes of Java over those of Objective-C?
10. What is the similarity between the class instances of Java and those of C#?
11. How many generic collections are present in the C# 2005 version?
12. What is a C# assembly?
13. Are Objective-C classes types?
14. What is a Ruby module?
15. What is the use of a C# destructor?
16. When are constructors implicitly called in Objective-C?
17. Why are the classes in Ruby called dynamic?
18. Why are the stacks in Ruby more flexible than those in C++ and Java?
19. What is the drawback of user-defined generic classes?
20. What is a dynamic link library?
21. Where are Java classes allocated?
22. Why are destructors not as frequently needed in Java as they are in C++?
23. What is a friend function? What is a friend class?
24. What is one reason Java does not have friend functions or friend classes?
25. Describe the fundamental differences between C# structs and its classes.
26. How is a struct object in C# created?
27. Explain the three reasons accessors to private types are better than making the types public.
28. What are the differences between a C++ struct and a C# struct?
29. What is the name of all Ruby constructors?
30. What is the fundamental difference between the classes of Ruby and those of C++ and Java?
31. How are instances of C++ template classes created?
32. Describe the two problems that appear in the construction of large programs that led to the development of encapsulation constructs.
33. What problems can occur using C to define abstract data types?
34. What is the advantage of using a C++ namespace?
35. What is a Java package, and what is its purpose?
36. Describe a .NET assembly.
37. What elements can appear in a Ruby module?
Problem Set

1. Some software engineers believe that abstraction is essential for software development. Do you agree? Support your answer.

2. Suppose someone designed a stack abstract data type in which the function top returned an access path (or pointer) rather than returning a copy of the top element. This is not a true data abstraction. Why? Give an example that illustrates the problem.

3. Write an analysis of the similarities of and differences between Java generic classes and C++ template classes.

4. Discuss the advantages of C# properties, relative to writing accessor methods in C++ or Java.

5. What would be the dangers if a language does not have the ability to hide information?

6. Why didn’t C++ eliminate the problems discussed in Problem 5?

7. What are the advantages and disadvantages of the Objective-C approach to syntactically distinguishing class methods from instance methods?

8. In what ways are the method calls in C++ more or less readable than those of Objective-C?

9. What are the arguments for and against the Objective-C design that method access cannot be restricted?

10. Why are destructors rarely used in Java but essential in C++?

11. What are the arguments for and against the Java policy on collection?

12. Describe a situation where a C++ class is preferable to a C++ struct.

13. Explain why naming encapsulations are important for developing large programs.

14. Describe the three ways a client can reference a name from a namespace in C++.

15. The namespace of the C# standard library, System, is not implicitly available to C# programs. Do you think this is a good idea? Defend your answer.

16. What are the advantages and disadvantages of the ability to change objects in Ruby?

17. Compare the encapsulation in C with that in C++.

Programming Exercises

1. Design an abstract data type for a matrix with integer elements in a language that you know, including operations for addition, subtraction, and matrix multiplication.
2. Design a queue abstract data type for float elements in a language that you know, including operations for enqueue, dequeue, and empty. The dequeue operation removes the element and returns its value.

3. Modify the C++ class for the abstract stack type shown in Section 11.4.2 to use a linked list representation and test it with the same code that appears in this chapter.

4. Modify the Java class for the abstract stack type shown in Section 11.4.4 to use a linked list representation and test it with the same code that appears in this chapter.

5. Write an abstract data type for complex numbers, including operations for addition, subtraction, multiplication, division, extraction of each of the parts of a complex number, and construction of a complex number from two floating-point constants, variables, or expressions. Use C++, Java, C#, or Ruby.

6. Write an abstract data type for queues whose elements store 10-character names. The queue elements must be dynamically allocated from the heap. Queue operations are enqueue, dequeue, and empty. Use either C++, Java, C#, or Ruby.

7. Write an abstract data type for a queue whose elements can be any primitive type. Use Java 5.0, C# 2005, or C++.

8. Write an abstract data type for a queue whose elements include both a 20-character string and an integer priority. This queue must have the following methods: enqueue, which takes a string and an integer as parameters; dequeue, which returns the string from the queue that has the highest priority; and empty. The queue is not to be maintained in priority order of its elements, so the dequeue operation must always search the whole queue.

9. A deque is a double-ended queue, with operations adding and removing elements from either end. Modify the solution to Programming Exercise 7 to implement a deque.

10. Write an abstract data type for rational numbers (a numerator and a denominator). Include a constructor and methods for getting the numerator, getting the denominator, addition, subtraction, multiplication, division, equality testing, and display. Use Java, C#, C++, or Ruby.
12.1 Introduction
12.2 Object-Oriented Programming
12.3 Design Issues for Object-Oriented Languages
12.4 Support for Object-Oriented Programming in Specific Languages
12.5 Implementation of Object-Oriented Constructs
12.6 Reflection
This chapter begins with a brief introduction to object-oriented programming, followed by an extended discussion of the primary design issues for inheritance and dynamic binding. Next, the support for object-oriented programming in Smalltalk, C++, Objective-C, Java, C#, and Ruby is discussed. Following this is a short overview of the implementation of dynamic bindings of method calls to methods in object-oriented languages. The last section discusses reflection.

12.1 Introduction

Languages that support object-oriented programming now are firmly entrenched in the mainstream. From COBOL to LISP, including virtually every language in between, dialects that support object-oriented programming have appeared. C++ and Objective-C support procedural and data-oriented programming, in addition to object-oriented programming. CLOS, an object-oriented version of LISP (Paepeke, 1993), also supports functional programming. Some of the newer languages that were designed to support object-oriented programming do not support other programming paradigms but still employ some of the basic imperative structures and have the appearance of the older imperative languages. Among these are Java and C#. Ruby is challenging to categorize: It is a pure object-oriented language in the sense that all data are objects, but it is a hybrid language in that one can use it for procedural programming. Finally, there is the pure object-oriented but somewhat unconventional language: Smalltalk. Smalltalk was the first language to offer complete support for object-oriented programming. The details of support for object-oriented programming vary widely among languages, and that is the primary topic of this chapter.

This chapter relies heavily on Chapter 11. It is in fact a continuation of that chapter. This relationship reflects the reality that object-oriented programming is, in essence, an application of the principle of abstraction to abstract data types. Specifically, in object-oriented programming, the commonality of a collection of similar abstract data types is factored out and put in a new type. The members of the collection inherit these common parts from that new type. This feature is inheritance, which is at the center of object-oriented programming and the languages that support it.

The other characterizing feature of object-oriented programming, dynamic binding of method calls to methods, is also extensively discussed in this chapter.

Although object-oriented programming is supported by some of the functional languages, for example, CLOS, OCaml, and F#, those languages are not discussed in this chapter.
12.2 Object-Oriented Programming

12.2.1 Introduction

The concept of object-oriented programming had its roots in SIMULA 67 but was not fully developed until the evolution of Smalltalk resulted in Smalltalk 80 (in 1980, of course). Indeed, some consider Smalltalk to be the base model for a purely object-oriented programming language. A language that is object-oriented must provide support for three key language features: abstract data types, inheritance, and dynamic binding of method calls to methods. Abstract data types were discussed in detail in Chapter 11, so this chapter focuses on inheritance and dynamic binding.

12.2.2 Inheritance

There has long been pressure on software developers to increase their productivity. This pressure has been intensified by the continuing reduction in the cost of computer hardware. By the middle to late 1980s, it became apparent to many software developers that one of the most promising opportunities for increased productivity in their profession was in software reuse. Abstract data types, with their encapsulation and access controls, are obvious candidates for reuse. The problem with the reuse of abstract data types is that, in nearly all cases, the features and capabilities of the existing type are not quite right for the new use. The old type requires at least some minor modifications. Such modifications can be difficult, because they require the person doing the modification to understand part, if not all, of the existing code. In many cases, the person doing the modification is not the program’s original author. Furthermore, the modifications often require changes to all client programs.

A second problem with programming with abstract data types is that the type definitions are all independent and are at the same level. This design often makes it impossible to organize a program to match the problem space being addressed by the program. In many cases, the underlying problem has categories of objects that are related, both as siblings (being similar to each other) and as parents and children (having a descendant relationship).

Inheritance offers a solution to both the modification problem posed by abstract data type reuse and the program organization problem. If a new abstract data type can inherit the data and functionality of some existing type, and is also allowed to modify some of those entities and add new entities, reuse is greatly facilitated without requiring changes to the reused abstract data type. Programmers can begin with an existing abstract data type and design a modified descendant of it to fit a new problem requirement. Furthermore,

1. Although SIMULA 67 had classes, the members defined inside them were not hidden from outside code.

2. This is similar to the functions in a C program, which are also independent and at a single level.
inheritance provides a framework for the definition of hierarchies of related classes that can reflect the descendant relationships in the problem space.

The abstract data types in object-oriented languages, following the lead of SIMULA 67, are usually called classes. As with instances of abstract data types, class instances are called objects. A class that is defined through inheritance from another class is a derived class, a subclass, or a child class. A class from which the new class is derived is its base class, superclass, or parent class. The subprograms that define the operations on objects of a class are called methods. The calls to methods are sometimes called messages. The entire collection of methods of a class is called the message protocol, or message interface, of the class. Computations in an object-oriented program are specified by messages sent from objects to other objects, or in some cases, to classes.

Methods are similar to subprograms. Both are collections of code that perform some computation. Both can take parameters and return results.

Passing a message is different from calling a subprogram. A subprogram typically processes data that is either passed to it by its caller as a parameter or is accessed nonlocally or globally. A message that is sent to an object is a request to execute one of its methods. At least some of the data on which the method is to operate is part of the object itself. Objects have methods that define processes the object can perform on itself. Because the objects are of abstract data types, these should be the only ways to manipulate data of the object. A subprogram defines a process that it can perform on any data sent to it (or made available nonlocally or globally).

As a simple example of inheritance, consider the following: Suppose we have a class named Vehicles, which has variables for year, color, and make. A natural specialization, or subclass, of this would be Truck, which could inherit the variables from Vehicle, but would add variables for hauling capacity and number of wheels. Figure 12.1 shows a simple diagram to indicate the relationship between the Vehicle class and the Truck class, in which the arrow points to the parent class.

There are several ways a derived class can differ from its parent. Following are the most common differences between a parent class and its subclasses:

1. The subclass can add variables and/or methods to those inherited from the parent class.
2. The subclass can modify the behavior of one or more of its inherited methods. A modified method has the same name, and often the same protocol, as the one of which it is a modification.

3. If a subclass does not differ from its parent, it obviously serves no purpose.

Figure 12.1
A simple example of inheritance

Vehicle

Truck
3. The parent class can define some of its variables or methods to have private access, which means they will not be visible in the subclass.

The new method is said to **override** the inherited method, which is then called an **overridden method**. The purpose of an overriding method is to provide an operation in the subclass that is similar to one in the parent class, but is customized for objects of the subclass. For example, a parent class, Bird, might have a draw method that draws a generic bird. A subclass of Bird named Waterfowl could override the draw method inherited from Bird to draw a generic waterfowl, perhaps a duck.

Classes can have two kinds of methods and two kinds of variables. The most commonly used methods and variables are called **instance methods** and **instance variables**. Every object of a class has its own set of instance variables, which store the object's state. The only difference between two objects of the same class is the state of their instance variables.

For example, a class for cars might have instance variables for color, make, model, and year. Instance methods operate only on the objects of the class. **Class variables** belong to the class, rather than its object, so there is only one copy for the class. For example, if we wanted to count the number of instances of a class, the counter could not be an instance variable—it would need to be a class variable. **Class methods** can perform operations on the class, and possibly also on the objects of the class. They can be called by prefixing their names with either the class name or a variable that references one of their instances. If a class defines a class method, that method can be called even if there are no instances of the class. A class method could be used to create an instance of the class.

If a new class is a subclass of a single parent class, then the derivation process is called **single inheritance**. If a class has more than one parent class, the process is called **multiple inheritance**. When a number of classes are related through single inheritance, their relationships to each other can be shown in a derivation tree. The class relationships in a multiple inheritance can be shown in a derivation graph. This is shown in Figure 12.5 in Section 12.4.2.2.

One disadvantage of inheritance as a means of increasing the possibility of reuse is that it creates dependencies among the classes in an inheritance hierarchy. This result works against one of the advantages of abstract data types, which is that they are independent of each other. Of course, not all abstract data types must be completely independent. But in general, the independence of abstract data types is one of their strongest positive characteristics. However, it may be difficult, if not impossible, to increase the reusability of abstract data types without creating dependencies among some of them. Furthermore, in many cases, the dependencies naturally mirror dependencies in the underlying problem space.

In Chapter 11 the access controls for variables and methods, together often called members, in a class are discussed. Private members are visible inside the class, while public members also are visible to clients of the class. Inheritance brings a new category of possible visibility, subclasses. Private members of a

---

4. This is not true in Ruby, which allows different objects of the same class to differ in other ways.
base class are not visible to subclasses, but public members are. The third level of accessibility, protected, allows members of a base class to be visible to subclasses, but not clients.

### 12.2.3 Dynamic Binding

The third essential characteristic (after abstract data types and inheritance) of object-oriented programming languages is a kind of polymorphism provided by the dynamic binding of messages to method definitions. This is sometimes called dynamic dispatch. Consider the following situation: There is a base class, `A`, that defines a method `draw` that draws some figure associated with the base class. A second class, `B`, is defined as a subclass of `A`. Objects of this new class also need a `draw` method that is like that provided by `A` but a bit different because the subclass objects are slightly different. So, the subclass overrides the inherited `draw` method. If a client of `A` and `B` has a variable that is a reference to class `A`'s objects, that reference also could point at class `B`'s objects, making it a polymorphic reference. If the method `draw`, which is defined in both classes, is called through the polymorphic reference, the run-time system must determine, during execution, which method should be called, `A`'s or `B`'s (by determining which type object is currently referenced by the reference). Figure 12.2 shows this situation.

Polymorphism is a natural part of any object-oriented language that is statically typed. In a sense, polymorphism makes a statically typed language a little bit dynamically typed, where the little bit is in some bindings of method calls to methods. The type of a polymorphic variable is indeed dynamic.

The approach just described is not the only way to design polymorphic references. One alternative, which is used in Objective-C, is described in Section 12.4.3.

One purpose of dynamic binding is to allow software systems to be more easily extended during both development and maintenance. Suppose we have

```java
public class A {
    ...
    draw( ) {...}
    ...
}  

public class B extends A {
    ...
    draw( ) {...}
    ...
}
```

---

5. Polymorphism is defined in Chapter 9.

6. Dynamic binding of method calls to methods is sometimes called dynamic polymorphism.
a catalog of used cars that is implemented as a `Car` class and a subclass for each car in the catalog. The subclasses contain an image of the car and specific information about the car. Users can browse the cars with a program that displays the images and information about each car as the user browses to it. The display of each car (and its information) includes a button that the user can click if he or she is interested in that particular car. After the user gets through the catalog, the system will print the images and information about the cars of interest to the user. One way to implement this system is to place a reference to each car (subclass of `Car`) of interest in a list that can store references to the base class, `Car`. When the user is ready, information about all of the cars of interest could be printed for the user to study and compare the cars in the list. The catalog of cars will of course change frequently. This will necessitate corresponding changes in the subclasses of `Car`. However, changes to the collection of subclasses will not require any other changes to the system.

In some cases, the design of an inheritance hierarchy results in one or more classes that are so high in the hierarchy that an instantiation of them would not make sense. For example, suppose a program defined a `Building` class and a collection of subclasses for specific types of buildings, for instance, `French_Gothic_Cathedrals`. It probably would not make sense to have an implemented `draw` method in `Building`. But because all of its descendant classes should have such methods, the protocol (but not the body) of that method is included in `Building`. Such a method is often called an abstract method (pure virtual method in C++). A class that includes at least one abstract method is called an abstract class (abstract base class in C++). Such a class usually cannot be instantiated, because some of its methods are declared but are not defined (they do not have bodies). Any subclass of an abstract class that is to be instantiated must provide implementations (definitions) of all of the inherited abstract methods.

### 12.3 Design Issues for Object-Oriented Languages

A number of issues must be considered when designing the programming language features to support inheritance and dynamic binding. Those that we consider most important are discussed in this section.

#### 12.3.1 The Exclusivity of Objects

A language designer who is totally committed to the object model of computation designs an object system that subsumes all other concepts of type. Everything, from a simple scalar integer to a complete software system, is an object in this mind-set. The advantage of this choice is the elegance and pure uniformity of the language and its use. The primary disadvantage is that simple operations must be done through the message-passing process, which often makes them slower than similar operations in an imperative model, where single machine
instructions implement such simple operations. In this purest model of object-oriented computation, all types are classes. There is no distinction between predefined and user-defined classes. In fact, all classes are treated the same way and all computation is accomplished through message passing.

One alternative to the exclusive use of objects that is common in imperative languages to which support for object-oriented programming has been added is the following: Retain the complete collection of types from the base imperative language and add the object typing model. This approach results in a larger language whose type structure can be confusing to new users of the language.

Another alternative to the exclusive use of objects is to have an imperative-style type structure for the primitive scalar types, but implement all structured types as objects. This choice provides the speed of operations on primitive values that is comparable to those expected in the imperative model.

12.3.2 Are Subclasses Subtypes?

If a language allows programs in which a variable of a class can be substituted for a variable of one of its ancestor classes in any situation, without causing type errors and without changing the behavior of the program, that language supports the principle of substitution. In such a language, if class B is derived from class A, then B has everything A has and the behavior of an object of class B, when used in place of an object of class A, is identical to that of an object of class A.\footnote{There is a fundamental theoretical problem with this last requirement: In general, it is not possible to determine if the behaviors of two programs are identical.} When this is true, B is a subtype of A. Although a subclass that is a subtype of its parent class must expose all of the members that are exposed by its parent class, the subclass can have members that are not in the parent class and still be a subtype.

The subtypes of Ada are examples of predefined subtypes. For example,

```plaintext
subtype Small_Int is Integer range -100..100;
```

Variables of `Small_Int` type have all of the operations of `Integer` variables but can store only a subset of the values possible in `Integer`. Furthermore, every `Small_Int` variable can be used anywhere an `Integer` variable can be used. That is, every `Small_Int` variable is, in a sense, an `Integer` variable.

The definition of subtype clearly disallows having public entities in the parent class that are not public in the subclass. So, the derivation process for subtypes must require that public entities of the parent class are inherited as public entities in the subclass.

Not all subcategories are subtypes and not all subtypes are subcategories. For example, a subclass cannot be a subtype if it changes the behavior of one of its overriding methods. Also, a class that is not a subclass of another class can be a subtype of that class by defining the same members, in terms of both types and
behavior. A subtype inherits interfaces and behavior, while a subclass inherits implementation, primarily to promote code reuse.

Most static-typed languages that support object-oriented programming are designed so that subclasses are subtypes, unless the programmer specifically designs a subclass that has behavior that differs from that of its parent class.

One obvious question is: Is the issue of whether subclasses are subtypes a theoretical or practical one? It is probably unusual to define a subclass whose overriding methods preserve the type protocols of their corresponding overridden methods but not their effects. So it is not a frequent practical issue. However, requiring all subclasses to be subtypes, if there were a reasonably simple way to enforce that, would place inheritance on a sounder theoretical base.

### 12.3.3 Single and Multiple Inheritance

Another simple design issue for object-oriented languages is: Does the language allow multiple inheritance (in addition to single inheritance)? Maybe it's not so simple. The purpose of multiple inheritance is to allow a new class to inherit from two or more classes.

Because multiple inheritance is sometimes highly useful, why would a language designer not include it? The reasons lie in two categories: complexity and efficiency. The additional complexity is illustrated by several problems. First, note that if a class has two unrelated parent classes and neither defines a name that is defined in the other, there is no problem. However, suppose a subclass named \( C \) inherits from both class \( A \) and class \( B \) and both \( A \) and \( B \) define an inheritable method named \( \text{display} \). If \( C \) needs to reference both versions of \( \text{display} \), how can that be done? This ambiguity problem is further complicated when the two parent classes both define identically named methods and one or both of them must be overridden in the subclass.

Another issue arises if both \( A \) and \( B \) are derived from a common parent, \( Z \), and \( C \) has both \( A \) and \( B \) as parent classes. This situation is called diamond or shared inheritance. In this case, both \( A \) and \( B \) should include \( Z \)'s inheritable variables. Suppose \( Z \) includes an inheritable variable named \( \text{sum} \). The question is whether \( C \) should inherit both versions of \( \text{sum} \) or just one, and if just one, which one? There may be programming situations in which just one of the two should be inherited, and others in which both should be inherited. A similar problem occurs when both \( A \) and \( B \) inherit a method from \( Z \) and both override that method. If a client of \( C \), which inherits both overriding methods, calls the method, which method is called, or are both supposed to be called. Diamond inheritance is shown in Figure 12.3.
The question of efficiency may be more perceived than real. In C++, for example, supporting multiple inheritance requires just one additional array access and one extra addition operation for each dynamically bound method call, at least with some machine architectures (Stroustrup, 1994, p. 270). Although this operation is required even if the program does not use multiple inheritance, it is a small additional cost.

The use of multiple inheritance can easily lead to complex program organizations. Many who have attempted to use multiple inheritance have found that designing the classes to be used as multiple parents is difficult. And the difficulties are not restricted to those created by the initial developer. A class might be used by another developer at some later date as one of the parents of a new class. Maintenance of systems that use multiple inheritance can be a more serious problem, for multiple inheritance leads to more complex dependencies among classes. It is not clear to some that the benefits of multiple inheritance are worth the added effort to design and maintain a system that uses it.

An interface is somewhat similar to an abstract class; its methods are declared but not defined. Interfaces cannot be instantiated. They are used as an alternative to multiple inheritance.8 Interfaces provide some of the benefits of multiple inheritance but have fewer disadvantages. For example, the problems of diamond inheritance are avoided when interfaces, rather than multiple inheritance, are used.

12.3.4 Allocation and Deallocation of Objects

There are two design questions concerning the allocation and deallocation of objects. The first of these is the place from which objects are allocated. If they behave like the abstract data types, then they can be allocated from anywhere. This means they could be allocated from the run-time stack or explicitly created on the heap with an operator or function, such as new. If they are all heap dynamic, there is the advantage of having a uniform method of creation and access through pointer or reference variables. This design simplifies the assignment operation for objects, making it in all cases only a pointer or reference value change. It also allows references to objects to be implicitly dereferenced, simplifying the access syntax.

If objects are stack dynamic, there is a potential problem with regard to subtypes. If class B is a child of class A and B is a subtype of A, then an object of B type can be assigned to a variable of A type. For example, if b1 is a variable of B type and a1 is a variable of A type, then

\[
a1 = b1;
\]

is a legal statement. If a1 and b1 are references to heap-dynamic objects, there is no problem—the assignment is a simple pointer assignment. However, if a1 and b1 are stack dynamic, then they are value variables and, if assigned the

---

8. Interfaces initially appeared in Java, whose designers recognized the difficulties that are raised by the use of multiple inheritance.
value of the object, must be copied to the space of the target object. If \( B \) adds a data field to what it inherited from \( A \), then \( a1 \) will not have sufficient space on the stack for all of \( b1 \). The excess will simply be truncated, which could be confusing to programmers who write or use the code. This truncation is called **object slicing**. The following example and Figure 12.4 illustrate the problem.

```cpp
class A {
    int x;
    ...;
};
class B : A {
    int y;
    ...;
}
```

The second question here is concerned with those cases where objects are allocated from the heap. The question is whether deallocation is implicit, explicit, or both. If deallocation is implicit, some implicit method of storage reclamation is required. If deallocation can be explicit, that raises the issue of whether dangling pointers or references can be created.

### 12.3.5 Dynamic and Static Binding

As discussed in Section 12.2.3, dynamic binding of messages to methods is an essential part of object-oriented programming. The question here is whether all bindings of messages to methods are dynamic. The alternative is to allow the user to specify whether a specific binding is to be dynamic or static. The advantage of this is that static bindings are faster. So, if a binding need not be dynamic, why pay the price?

### 12.3.6 Nested Classes

One of the primary motivations for nesting class definitions is information hiding. If a new class is needed by only one class, there is no reason to define it so it can be seen by other classes. In this situation, the new class can be nested
inside the class that uses it. In some cases, the new class is nested inside a sub-program, rather than directly in another class.

The class in which the new class is nested is called the nesting class. The most obvious design issues associated with class nesting are related to visibility. Specifically, one issue is: Which of the members of the nesting class are visible in the nested class? The other important issue is the opposite: Which of the members of the nested class are visible in the nesting class?

### 12.3.7 Initialization of Objects

The initialization issue is whether and how objects are initialized to values when they are created. This is more complicated than may be first thought. One question is whether objects must be initialized manually or through some implicit mechanism. When an object of a subclass is created, is the associated initialization of the inherited parent class member implicit or must the programmer explicitly deal with it?

### 12.4 Support for Object-Oriented Programming in Specific Languages

#### 12.4.1 Smalltalk

Many think of Smalltalk as the definitive object-oriented programming language. It was the first language to include complete support for that paradigm. Therefore, it is natural to begin a survey of language support for object-oriented programming with Smalltalk.

##### 12.4.1.1 General Characteristics

In Smalltalk, the concept of an object is truly universal. Virtually everything, from things as simple as the integer constant 2 to a complex file-handling system, is an object. As objects, they are treated uniformly. They all have local memory, inherent processing ability, the capability to communicate with other objects, and the possibility of inheriting methods and instance variables from ancestors. Classes cannot be nested in Smalltalk.

All computation is through messages, even a simple arithmetic operation. For example, the expression \( x + 7 \) is implemented as sending the \(+\) message to \( x \) (to enact the \(+\) method), sending 7 as the parameter. This operation returns a new numeric object with the result of the addition.

Replies to messages have the form of objects and are used to return requested or computed information or only to confirm that the requested service has been completed.

All Smalltalk objects are allocated from the heap and are referenced through reference variables, which are implicitly dereferenced. There is no
explicit deallocation statement or operation. So, all deallocation is implicit, using a garbage collection process for storage reclamation.

In Smalltalk, constructors must be explicitly called when an object is created. A class can have multiple constructors, but each must have a unique name. Smalltalk classes cannot be nested in other classes.

Unlike hybrid languages such as C++ and Objective-C, Smalltalk was designed for just one software development paradigm—object oriented. Furthermore, it adopts none of the appearance of the imperative languages. Its purity of purpose is reflected in its simple elegance and uniformity of design.

There is an example Smalltalk program in Chapter 2.

### 12.4.1.2 Inheritance

A Smalltalk subclass inherits all of the members of its superclass. The subclass can also have its own instance variables, which must have names that are distinct from the variable names in its ancestor classes. Finally, the subclass can define new methods and redefine methods that already exist in an ancestor class. When a subclass has a method whose name and protocol are the same as an ancestor class, the subclass method hides that of the ancestor class. Access to such a hidden method is provided by prefixing the message with the pseudovariable `super`. The prefix causes the method search to begin in the superclass rather than locally.

Because members in a parent class cannot be hidden from subclasses, subclasses can be and usually are subtypes.

Smalltalk does not support multiple inheritance.

### 12.4.1.3 Dynamic Binding

The dynamic binding of messages to methods in Smalltalk operates as follows: A message to an object causes a search of the class to which the object belongs for a corresponding method. If the search fails, it is continued in the superclass of that class, and so forth, up to the system class, `Object`, which has no superclass. `Object` is the root of the class derivation tree on which every class is a node. If no method is found anywhere in that chain, an error occurs. It is important to remember that this method search is dynamic—it takes place when the message is sent. Smalltalk does not, under any circumstances, bind messages to methods statically.

The only type checking in Smalltalk is dynamic, and the only type error occurs when a message is sent to an object that has no matching method, either locally or through inheritance. This is a different concept of type checking than that of most other languages. Smalltalk type checking has the simple goal of ensuring that a message matches some method.

Smalltalk variables are not typed; any name can be bound to any object. As a direct result, Smalltalk supports dynamic polymorphism. All Smalltalk code is generic in the sense that the types of the variables are irrelevant, as long as they are consistent. The meaning of an operation (method or operator) on a variable is determined by the class of the object to which the variable is currently bound.
The point of this discussion is that as long as the objects referenced in an expression have methods for the messages of the expression, the types of the objects are irrelevant. This means that no code is tied to a particular type.

### 12.4.1.4 Evaluation of Smalltalk

Smalltalk is a small language, although the Smalltalk system is large. The syntax of the language is simple and highly regular. It is a good example of the power that can be provided by a small language if that language is built around a simple but powerful concept. In the case of Smalltalk, that concept is that all programming can be done employing only a class hierarchy built using inheritance, objects, and message passing.

In comparison with conventional compiled imperative-language programs, equivalent Smalltalk programs are significantly slower. Although it is theoretically interesting that array indexing and loops can be provided within the message-passing model, efficiency is an important factor in the evaluation of programming languages. Therefore, efficiency will clearly be an issue in most discussions of the practical applicability of Smalltalk.

Smalltalk’s dynamic binding allows type errors to go undetected until run time. A program can be written that includes messages to nonexistent methods and it will not be detected until the messages are sent, which causes a great deal more error repair later in the development than would occur in a statically-typed language. However, in practice type errors are not a serious problem with Smalltalk programs.

Overall, the design of Smalltalk consistently came down on the side of language elegance and strict adherence to the principles of object-oriented programming support, often without regard for practical matters, in particular execution efficiency. This is most obvious in the exclusive use of objects and the typeless variables.

The Smalltalk user interface has had an important impact on computing: The integrated use of windows, mouse-pointing devices, and pop-up and pull-down menus, all of which first appeared in Smalltalk, dominate contemporary software systems.

Perhaps the greatest impact of Smalltalk is the advancement of object-oriented programming, now the most widely used design and coding methodology.

### 12.4.2 C++

Chapter 2 describes how C++ evolved from C and SIMULA 67, with the design goal of support for object-oriented programming while retaining nearly complete backward compatibility with C. C++ classes, as they are used to support abstract data types, are discussed in Chapter 11. C++ support for the other essentials of object-oriented programming is explored in this section. The whole collection of details of C++ classes, inheritance, and dynamic binding is large and complex. This section discusses only the most important among these topics, specifically, those directly related to the design issues described in Section 12.3.
C++ was the first widely used object-oriented programming language and is still among the most popular. So, naturally, it is the one with which other languages are often compared. For both of these reasons, our coverage of C++ here is more detailed than that of the other example languages discussed in this chapter.

12.4.2.1 General Characteristics

To maintain backward compatibility with C, C++ retains the type system of C and adds classes to it. Therefore, C++ has both traditional imperative-language types and the class structure of an object-oriented language. It supports methods, as well as functions that are not related to specific classes. This makes it a hybrid language, supporting both procedural programming and object-oriented programming.

The objects of C++ can be static, stack dynamic, or heap dynamic. Explicit deallocation using the \texttt{delete} operator is required for heap-dynamic objects, because C++ does not include implicit storage reclamation.

Many class definitions include a destructor method, which is implicitly called when an object of the class ceases to exist. The destructor is used to deallocate heap-allocated memory that is referenced by data members. It may also be used to record part or all of the state of the object just before it dies, usually for debugging purposes.

12.4.2.2 Inheritance

A C++ class can be derived from an existing class, which is then its parent, or base, class. Unlike Smalltalk and most other languages that support object-oriented programming, a C++ class can also be stand-alone, without a superclass. In the definition of a derived class, the name of the derived class has the name of the base class attached with a colon (:), as in the following syntactic form:

\begin{verbatim}
class derived_class_name : base_class_name { ... }
\end{verbatim}

The data defined in a class definition are called \textit{data members} of that class, and the functions defined in a class definition are called \textit{member functions} of that class (member functions in other languages are usually called methods). Some or all of the members of the base class may be inherited by the derived class, which can also add new members and modify inherited member functions.

All C++ objects must be initialized before they are used. Therefore, all C++ classes include at least one constructor method that initializes the data members of the new object. Constructor methods are implicitly called when an object is created. If any of the data members are pointers to heap-allocated data, the constructor allocates that storage.

If a class is derived from another class, the inherited data members must be initialized when the derived class object is created. To do this, the base class constructor is implicitly called. When initialization data must be furnished to
On Paradigms and Better Programming

BJARNE STROUSTRUP

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(year of interview: 2002)

PROGRAMMING PARADIGMS

Your thoughts on the object-oriented paradigm: Its pluses and minuses. Let me first say what I mean by OOP—too many people think that “object-oriented” is simply a synonym for “good.” If so, there would be no need for other paradigms. The key to OO is the use of class hierarchies providing polymorphic behavior through some rough equivalent of virtual functions. For proper OO, it is important to avoid directly accessing the data in such a hierarchy and to use only a well-designed functional interface.

In addition to its well-documented strengths, object-oriented programming also has obvious weaknesses. In particular, not every concept naturally fits into a class hierarchy, and the mechanisms supporting object-oriented programming can impose significant overheads compared to alternatives. For many simple abstractions, classes that do not rely on hierarchies and run-time binding provide a simpler and more efficient alternative. Furthermore, where no run-time resolution is needed, generic programming relying on (compile-time) parametric polymorphism is a better behaved and more efficient approach.

So, C++: Is it O0 or other? C++ supports several paradigms—including OOP, generic programming, and procedural programming—and combinations of these paradigms define multiparadigm programming as supporting more than one programming style (“paradigm”) and combinations of those styles.

Do you have a mini-example of multiparadigm programming? Consider this variant of the classic “collection of shapes” examples (originating from the early days of the first language to support object-oriented programming: Simula 67):

```cpp
void draw_all(const vector<Shape*>& vs)
{
    for (int i = 0; i<vs.size(); ++i)
        vs[i]->draw();
}
```

Here, I use the generic container vector together with the polymorphic type Shape. The vector provides static type safety and optimal run-time performance. The Shape provides the ability to handle
a `Shape` (i.e., any object of a class derived from `Shape`) without recompilation.

We can easily generalize this to any container that meets the C++ standard library requirements:

```cpp
template<class C>
void draw_all(const C& c) {
  typedef typename C::const_iterator CI;
  for (CI p = c.begin(); p!=c.end(); ++p)
    (*p)->draw();
}
```

Using iterators allows us to apply this `draw_all()` to containers that do not support subscripts, such as a standard library list:

```cpp
vector<Shape*> vs;
list<Shape*> ls;
// . . .
draw_all(vs);
draw_all(ls);
```

We can even generalize this further to handle any sequence of elements defined by a pair of iterators:

```cpp
template<class Iterator>
void draw_all(Iterator b, Iterator e) {
  for_each(b,e,mem_fun(&Shape::draw));
}
```

To simplify the implementation, I used the standard library algorithm `for_each`.

We might call this last version of `draw_all()` for a standard library list and an array:

```cpp
list<Shape*> ls;
Shape* as[100];
// . . .
draw_all(ls.begin(),ls.end());
draw_all(as,as+100);
```

**SELECTING THE “RIGHT” LANGUAGE FOR THE JOB**

How useful is it to have this background in numerous paradigms? Or would it be better to invest time in becoming even more familiar with OO languages rather than learning these other paradigms? It is essential for anyone who wants to be considered a professional in the areas of software to know several languages and several programming paradigms. Currently, C++ is the best language for multiparadigm programming and a good language for learning various forms of programming. However, it’s not a good idea to know just C++, let alone to know just a single-paradigm language. That would be a bit like being colorblind or monoglot: You would hardly know what you were missing. Much of the inspiration to good programming comes from having learned and appreciated several programming styles and seen how they can be used in different languages.

Furthermore, I consider programming of any non-trivial program a job for professionals with a solid and broad education, rather than for people with a hurried and narrow “training.”
the base class constructor, it is given in the call to the derived object constructor. In general, this is done with the following construct:

```
subclass (subclass parameters) : base_class (superclass parameters) {
    ...
}
```

If no constructor is included by the developer in a class definition, the compiler includes a trivial constructor. This default constructor calls the constructor of the base class, if there is a base class.

Class members can be private, protected, or public. Private members are accessible only by member functions and friends of the class. Standalone functions, member functions, and classes can be declared to be friends of a class and thereby be given access to its private members. Public members are visible everywhere. Protected members are like private members, except in derived classes, whose access is described next. Derived classes can modify accessibility for their inherited members. The complete syntactic form of a derived class is as follows:

```
class derived_class_name : derivation_mode base_class_name
    { data member and member function declarations };
```

The derivation_mode can be either public or private.\(^9\) (Do not confuse public and private derivation with public and private members.) The public and protected members of a base class are also public and protected, respectively, in a public-derived class. In a private-derived class, both the public and protected members of the base class are private. So, in a class hierarchy, a private-derived class cuts off access to all members of all ancestor classes to all successor classes. Private members of a base class are inherited by a derived class, but they are not visible to the members of that derived class and are therefore of no use there. Private derivations provide the possibility that a subclass can have members with different access than the same members in the parent class. Consider the following example:

```
class base_class {
    private:
        int a;
        float x;
    protected:
        int b;
        float y;
    public:
        int c;
        float z;
};

class subclass_1 : public base_class { ...};
class subclass_2 : private base_class { ...};
```

\(^9\) It can also be protected, but that option is not discussed here.
In subclass_1, b and y are protected, and c and z are public. In subclass_2, b, y, c, and z are private. No derived class of subclass_2 can have members with access to any member of base_class. The data members a and x in base_class are not accessible in either subclass_1 or subclass_2.

Note that private-derived subclasses cannot be subtypes. For example, if the base class has a public data member, under private derivation that data member would be private in the subclass. Therefore, if an object of the subclass were substituted for an object of the base class, accesses to that data member would be illegal on the subclass object. However, public-derived subclasses can be and usually are subtypes.

Under private class derivation, no member of the parent class is implicitly visible to the instances of the derived class. Any member that must be made visible must be reexported in the derived class. This reexportation in effect exempts a member from being hidden even though the derivation was private. For example, consider the following class definition:

```cpp
class subclass_3 : private base_class {
    base_class :: c;
    ...
}
```

Now, instances of subclass_3 can access c. As far as c is concerned, it is as if the derivation had been public. The double colon (::) in this class definition is a scope resolution operator. It specifies the class where its following entity is defined.

The example in the following paragraphs illustrates the purpose and use of private derivation.

Consider the following example of C++ inheritance, in which a general linked-list class is defined and then used to define two useful subclasses:

```cpp
class single_linked_list {
    private:
        class node {
            public:
                node *link;
                int contents;
            }
        node *head;
    public:
        single_linked_list() {head = 0};
        void insert_at_head(int);
        void insert_at_tail(int);
        int remove_at_head();
        int empty();
    }
}
```

The nested class, node, defines a cell of the linked list to consist of an integer variable and a pointer to a node object. The node class is in the private clause,
which hides it from all other classes. Its members are public, however, so they are visible to the nesting class, `single_linked_list`. If they were private, `node` would need to declare the nesting class to be a friend to make them visible in the nesting class. Note that nested classes have no special access to members of the nesting class. Only static data members of the nesting class are visible to methods of the nested class.\textsuperscript{10}

The enclosing class, `single_linked_list`, has just a single data member, a pointer to act as the list’s header. It contains a constructor function, which sets `head` to the null pointer value. The four member functions allow nodes to be inserted at either end of a list object, nodes to be removed from one end of a list, and lists to be tested for empty.

The following definitions provide stack and queue classes, both based on the `single_linked_list` class:

```cpp
class stack : public single_linked_list {
   public:
      stack() {}
      void push(int value) {
         insert_at_head(value);
      }
      int pop() {
         return remove_at_head();
      }
};

class queue : public single_linked_list {
   public:
      queue() {}
      void enqueue(int value) {
         insert_at_tail(value);
      }
      int dequeue() {
         remove_at_head();
      }
};
```

Note that objects of both the `stack` and `queue` subclasses can access the `empty` function defined in the base class, `single_linked_list` (because it is a public derivation). Both subclasses define constructor functions that do nothing. When an object of a subclass is created, the proper constructor in the subclass is implicitly called. Then, any applicable constructor in the base class is called. So, in our example, when an object of type `stack` is created, the constructor in `single_linked_list` is called, which does the necessary initialization. Then the `stack` constructor is called, which does nothing.

The classes `stack` and `queue` both suffer from the same serious problem: Clients of both can access all of the public members of the parent class,\textsuperscript{10}

\textsuperscript{10}. A class can also be defined in a method of a nesting class. The scope rules of such classes are the same as those for classes nested directly in other classes, even for the local variables declared in the method in which they are defined.
single-linked-list. A client of a stack object could call `insert_at_tail`, thereby destroying the integrity of its stack. Likewise, a client of a queue object could call `insert_at_head`. These unwanted accesses are allowed because both stack and queue are subtypes of single-linked-list. Public derivation is used when one wants the subclass to inherit the entire interface of the base class. The alternative is to use a derivation in which the subclass inherits only the implementation of the base class. Our two example derived classes can be written to make them not subtypes of their parent class by using private, rather than public, derivation. Then, both will also need to reexport empty, because it will become hidden to their instances. This situation illustrates the motivation for the private-derivation option. The new definitions of the stack and queue types, named `stack_2` and `queue_2`, are shown in the following:

```cpp
class stack_2 : private single_linked_list {
    public:
        stack_2() {}
        void push(int value) {
            single_linked_list :: insert_at_head(value);
        }
        int pop() {
            return single_linked_list :: remove_at_head();
        }
        single_linked_list:: empty();
};
class queue_2 : private single_linked_list {
    public:
        queue_2() {}
        void enqueue(int value) {
            single_linked_list :: insert_at_tail(value);
        }
        int dequeue() {
            single_linked_list :: remove_at_head();
        }
        single_linked_list:: empty();
};
```

These two classes use reexportation to allow access to base class methods for clients. This was not necessary when public derivation was used.

The two versions of stack and queue illustrate the difference between subtypes and derived types that are not subtypes. The linked list is a generalization of both stacks and queues, because both can be implemented as linked lists. So, it is natural to inherit from a linked-list class to define stack and queue classes. However, neither is a subtype of the linked-list class, because both make the public members of the parent class private, which makes them inaccessible to clients.

One of the reasons friends are necessary is that sometimes a subprogram must be written that can access the members of two different classes. For

11. They would not be subtypes because the public members of the parent class can be seen in a client, but not in a client of the subclass, where those members are private.
example, suppose a program uses a class for vectors and one for matrices, and a subprogram is needed to multiply a matrix object by a vector object. In C++, the multiply function can be made a friend of both classes.

As previously stated, C++ provides multiple inheritance. As an example, suppose we wanted a class for drawing that needed the behavior of a class written for drawing figures and the methods of the new class needed to run in a separate thread. We might define the following:

```c++
class Thread { . . . };
class Drawing { . . . };
class DrawThread : public Thread, public Drawing { . . . };
```

Class `DrawThread` inherits all of the members of both `Thread` and `Drawing`. If both `Thread` and `Drawing` happen to include members with the same name, they can be unambiguously referenced in objects of class `DrawThread` by using the scope resolution operator (`::`). This example of multiple inheritance is shown in Figure 12.5.

Some issues with the C++ implementation of multiple inheritance are discussed in Section 12.5.

Overriding methods\(^\text{12}\) in C++ must have exactly the same parameter profile as the overridden method. If there is any difference in the parameter profiles, the method in the subclass is considered a new method that is unrelated to the method with the same name in the ancestor class. The return type of the overriding method either must be the same as that of the overridden method or must be a publicly derived type of the return type of the overridden method.

### 12.4.2.3 Dynamic Binding

All of the member functions we have defined thus far are statically bound; that is, a call to one of them is statically bound to a function definition. A C++ object could be manipulated through a value variable, rather than a pointer or a reference. (Such an object would be static or stack dynamic.) However, in that

---

12. Recall that an overriding method is one that is defined in the derived class to replace a virtual method inherited from an ancestor class. Calls to an overriding method must be dynamically bound.
case, the object's type is known and static, so dynamic binding is not needed. On the other hand, a pointer variable that has the type of a base class can be used to point to any heap-dynamic object of any class publicly derived from that base class, making it a polymorphic variable. Publicly derived subclasses are subtypes if none of the members of the base class are private. Privately derived subclasses are never subtypes. A pointer to a base class cannot be used to reference a method in a subclass that is not a subtype.

C++ does not allow value variables (as opposed to pointers or references) to be polymorphic. When a polymorphic variable is used to call a member function overridden in one of the derived classes, the call must be dynamically bound to the correct member function definition. Member functions that must be dynamically bound must be declared to be virtual functions by preceding their headers with the reserved word \texttt{virtual}, which can appear only in a class body.

Consider the situation of having a base class named \texttt{Shape}, along with a collection of derived classes for different kinds of shapes, such as circles, rectangles, and so forth. If these shapes need to be displayed, then the displaying member function, \texttt{draw}, must be unique for each descendant, or kind of shape. These versions of \texttt{draw} must be defined to be virtual. When a call to \texttt{draw} is made with a pointer to the base class of the derived classes, that call must be dynamically bound to the member function of the correct derived class. The following example has the skeletal definitions for the example situation just described:

class Shape { 
  public: 
    virtual void draw() = 0; 
  . . .
}; 
class Circle : public Shape { 
  public: 
    void draw() { . . . } 
  . . .
}; 
class Rectangle : public Shape { 
  public: 
    void draw() { . . . } 
  . . .
}; 
class Square : public Rectangle { 
  public: 
    void draw() { . . . } 
  . . .
};

Given these definitions, the following code has examples of both statically and dynamically bound calls:

Square* sq = \texttt{new} Square; 
Rectangle* rect = \texttt{new} Rectangle; 
Shape* ptr_shape;
ptr_shape = sq;       // Now ptr_shape points to a
// Square object
ptr_shape->draw();    // Dynamically bound to the draw
// in the Square class
rect->draw();         // Statically bound to the draw
// in the Rectangle class

This situation is shown in Figure 12.6.

Notice that the draw function in the definition of the base class shape is set to 0. This peculiar syntax is used to indicate that this member function is a **pure virtual function**, meaning that it has no body and it cannot be called. It must be redefined in derived classes if they call the function. The purpose of a pure virtual function is to provide the interface of a function without giving any of its implementation. Pure virtual functions are usually defined when an actual member ...

---

**Figure 12.6**  
Dynamic binding

---

**Class Hierarchy**

```
Shape

virtual void draw() = 0
```

```
Circle

void draw() { ... }
```

```
Rectangle

void draw() { ... }
```

```
Square

void draw() { ... }
```

---

**Types**

```
Shape*

ptr_shape
```

```
Square*

sq
```

---

**Pointers**

```
Rectangle*

rect
```

---

**Bindings**

```
Square

void draw() { ... }
```

---

**Objects**

```
Rectangle

void draw() { ... }
```
function in the base class would not be useful. Recall that in Section 12.2.3, a base class `Building` was discussed, and each subclass described some particular kind of building. Each subclass had a `draw` method but none of these would be useful in the base class. So, `draw` would be a pure virtual function in the `Building` class.

Any class that includes a pure virtual function is an abstract class. In C++, an abstract class is not marked with a reserved word. An abstract class can include completely defined methods. Because of the presence of one or more virtual functions, it is illegal to instantiate an abstract class. In a strict sense, an abstract class is one that is used only to represent the characteristics of a type. C++ provides abstract classes to model these truly abstract classes. If a subclass of an abstract class does not redefine a pure virtual function of its parent class, that function remains as a pure virtual function in the subclass and the subclass is also an abstract class.

Abstract classes and inheritance together support a powerful technique for software development. They allow types to be hierarchically defined so that related types can be subclasses of truly abstract types that define their common abstract characteristics.

Dynamic binding allows the code that uses members like `draw` to be written before all or even any of the versions of `draw` are written. New derived classes could be added years later, without requiring any change to the code that uses such dynamically bound members. This is a highly useful feature of object-oriented languages.

Reference assignments for stack-dynamic objects are different from pointer assignments for heap-dynamic objects. For example, consider the following code, which uses the same class hierarchy as the last example:

```c++
Square sq; // Allocate a Square object on the stack
Rectangle rect; // Allocate a Rectangle object on
// the stack
rect = sq; // Copies the data member values from
// the Square object
rect.draw(); // Calls the draw from the Rectangle
// object
```

In the assignment `rect = sq`, the member data from the object referenced by `sq` would be assigned to the data members of the object referenced by `rect`, but `rect` would still reference the `Rectangle` object. Therefore, the call to `draw` through the object referenced by `rect` would be that of the `Rectangle` class. If `rect` and `sq` were pointers to heap-dynamic objects, the same assignment would be a pointer assignment, which would make `rect` point to the `Square` object, and a call to `draw` through `rect` would be bound dynamically to the `draw` in the `Square` object.

### 12.4.2.4 Evaluation

It is natural to compare the object-oriented features of C++ with those of Smalltalk. The inheritance of C++ is more intricate than that of Smalltalk in terms of access control. By using both the access controls within the class
definition and the derivation access controls, and also the possibility of friend functions and friend classes, the C++ programmer has highly detailed control over access to class members. Although C++ provides multiple inheritance and Smalltalk does not, there are many who feel that is not an advantage for C++. The downsides of multiple inheritance weigh heavily against its value. In fact, C++ is the only language discussed in this chapter that supports multiple inheritance. On the other hand, languages that provide alternatives to multiple inheritance, such as Objective-C, Java, and C#, clearly have an advantage over Smalltalk in that area.

In C++, the programmer can specify whether static binding or dynamic binding is to be used. Because static binding is faster, this is an advantage for those situations where dynamic binding is not necessary. Furthermore, even the dynamic binding in C++ is fast when compared with that of Smalltalk. Binding a virtual member function call in C++ to a function definition has a fixed cost, regardless of how distant in the inheritance hierarchy the definition appears. Calls to virtual functions require only five more memory references than statically bound calls (Stroustrup, 1988). In Smalltalk, however, messages are always dynamically bound to methods, and the farther away in the inheritance hierarchy the correct method is, the longer it takes. The disadvantage of allowing the user to decide which bindings are static and which are dynamic is that the original design must include these decisions, which may have to be changed later.

The static type checking of C++ is an advantage over Smalltalk, in which all type checking is dynamic. A Smalltalk program can be written with messages to nonexistent methods, which are not discovered until the program is executed. A C++ compiler finds such errors. Compiler-detected errors are less expensive to repair than those found in testing.

Smalltalk is essentially typeless, meaning that all code is effectively generic. This provides a great deal of flexibility, but static type checking is sacrificed. C++ provides generic classes through its template facility (as described in Chapter 11), which retains the benefits of static type checking.

The primary advantage of Smalltalk lies in the elegance and simplicity of the language, which results from the single philosophy of its design. It is purely and completely devoted to the object-oriented paradigm, devoid of compromises necessitated by the whims of an entrenched user base. C++, on the other hand, is a large and complex language with no single philosophy as its foundation, except to support object-oriented programming and include the C user base. One of its most significant goals was to preserve the efficiency and flavor of C while providing the advantages of object-oriented programming. Some people feel that the features of this language do not always fit well together and that at least some of the complexity is unnecessary.

According to Chambers and Ungar (1991), Smalltalk ran a particular set of small C-style benchmarks at only 10 percent of the speed of optimized C. C++ programs require only slightly more time than equivalent C programs (Stroustrup, 1988). Given the great efficiency gap between Smalltalk and C++, it is little wonder that the commercial use of C++ is far more widespread than that
of Smalltalk. There are other factors in this difference, but efficiency is clearly a strong argument in favor of C++. Of course, all of the compiled languages that support object-oriented programming run approximately 10 times faster than Smalltalk.

12.4.3 Objective-C

We discuss the support for object-oriented programming in Objective-C relative to that of C++. These two languages were designed at approximately the same time. Both add support for object-oriented programming to the C language. In appearance, the largest difference is in the syntax of method calls, which in C++ are closely related to the function calls of C, whereas in Objective-C they are more similar to the method calls of Smalltalk.

12.4.3.1 General Characteristics

Objective-C, like C#, has both primitive types and objects. Recall (from Chapter 11) that a class definition consists of two parts: interface and implementation. These two parts are often placed in separate files, the interface file using the .h name extension and the implementation using the .m name extension. When the interface is in a separate file, the implementation file begins with the following:

```c
#import "interface_file.h"
```

Instance variables are declared in a brace-delimited block following the header of the interface section. Objective-C does not support class variables directly. However, a static global variable that is defined in the implementation file can be used as a class variable.

The implementation section of a class contains definitions of the methods declared in the corresponding interface section.

Objective-C does not allow classes to be nested.

12.4.3.2 Inheritance

Objective-C supports only single inheritance. Every class must have a parent class, except the predefined root class named NSObject. One reason to have a single root class is that there are some operations that are universally needed. Among these are the class methods alloc and init. The parent class of a new class is declared in the interface directive after the colon that is attached to the name of the class being defined, as in the following:

```c
@interface myNewClass: NSObject {
```

Of course, all of the protected and public data members of the parent class are inherited by the subclass. New methods and instance variables can be added
to the subclass. Recall that although access to methods can be controlled by specifiers in C++, in Objective-C all methods are public, and that cannot be changed. A method that is defined in the subclass and has the same name, same return type, and same number and types of parameters overrides the inherited method. The overridden method can be called in another method of the subclass through super, a reference to the parent object. There is no way to prevent the overriding of an inherited method. Because all public members of a base class are also public in subclasses derived from that base class, all subclasses can be subtypes.

As in Smalltalk, in Objective-C any method name can be called on any object. If the run-time system discovers that the object has no such method (with the proper protocol), an error occurs.

Objective-C does not support the private and protected derivations of C++.

As in other languages that support object-oriented programming, the constructor of an instance of a subclass should always call the constructor of the parent class before doing anything else. If the name of the parent class constructor is init, this is done with the following statement:

```
[super init];
```

Objective-C includes two ways to extend a class besides subclassing: categories and protocols. A collection of methods can be added to a class with a construct called a category. A category is a secondary interface of a class that contains declarations of methods. No new instance variables can be included in the secondary interface. The syntactic form of such an interface is exemplified by the following:

```c
#import "Stack.h"
@interface Stack (StackExtend)
-(int) secondFromTop;
-(void) full;
@end
```

The name of this category is StackExtend. The original interface is accessible because it is imported, so the parent class need not be mentioned. The new methods are mixed into the methods of the original interface. Consequently, categories are sometimes called mixins. Mixins are sometimes used to add certain functionalities to different classes. And, of course, the class still has a normal superclass from which it inherits members. So, mixins provide some of the benefits of multiple inheritance, without the naming collisions that could occur if modules did not require module names on their functions. Of course, a category must also have an implementation section, which includes the name of the category in parentheses after the class name on the implementation directive, as in the following:

```c
@implementation Stack (StackExtend)
```

The implementation need not implement all of the methods in the category.
There is another way to provide some of the benefits of multiple inheritance in Objective-C, protocols. Although Objective-C does not provide abstract classes, as in C++, protocols are related to them. A *protocol* is a list of method declarations. The syntax of a protocol is exemplified with the following:

```objective-c
@protocol MatrixOps
-(Matrix *) add: (Matrix *) mat;
-(Matrix *) subtract: (Matrix *) mat;
@end
```

In this example, *MatrixOps* is the name of the protocol. The *add* and *subtract* methods must be implemented by a class that uses the protocol. This use is called *implementing* or *adopting* the protocol. The optional part specifies that the *multiply* method may or may not be implemented by an adopting class.

A class that adopts a protocol lists the name of the protocol in angle brackets after the name of the class on the interface directive, as in the following:

```objective-c
@interface MyClass: NSObject <YourProtocol>
@end
```

### 12.4.3.3 Dynamic Binding

In Objective-C, polymorphism is implemented in a way that differs from the way it is done in most other common programming languages. A polymorphic variable is created by declaring it to be of type *id*. Such a variable can reference any object. The run-time system keeps track of the class of the object to which an *id* type variable refers. If a call to a method is made through such a variable, the call is dynamically bound to the correct method, assuming one exists.

For example, suppose that a program has classes defined named *Circle* and *Square* and both have methods named *draw*. Consider the following skeletal code:

```objective-c
// Create the objects
Circle *myCircle = [[Circle alloc] init];
Square *mySquare = [[Square alloc] init];

// Initialize the objects
[myCircle setCircumference: 5];
[mySquare setSide: 5];

// Create the id variable
id shapeRef;

// Set the id to reference the circle and draw it
shapeRef = myCircle;
[shapeRef draw];
```
/ Set the id to reference the square
shapeRef = mySquare;
[shapeRef draw];

This code first draws the circle and then the square, with both draw methods called through the shapeRef object reference.

12.4.3.4 Evaluation

The support for object-oriented programming in Objective-C is adequate, although there are a few minor deficiencies. There is no way to prevent over-riding of an inherited method. Support for polymorphism with its id data type is overkill, for it allows variables to reference any object, rather than just those in an inheritance line. Although there is no direct support for multiple inheritance, the language includes a form of a mixin, categories, which provide some of the capabilities of multiple inheritance, without all of its disadvantages. Categories allow a collection of behaviors to be added to any class. Protocols provide the capabilities of interfaces, such as those in Java, which also provide some of the capabilities of multiple inheritance.

12.4.4 Java

Because Java’s design of classes, inheritance, and methods is similar to that of C++, in this section we focus only on those areas in which Java differs from C++.

12.4.4.1 General Characteristics

As with C++, Java supports both objects and nonobject data. However, in Java, only values of the primitive scalar types (Boolean, character, and the numeric types) are not objects. Java’s enumerations and arrays are objects. The reason Java has nonobjects is efficiency.

In Java 5.0+, primitive values are implicitly coerced when they are put in object context. This coercion converts the primitive value to an object of the wrapper class of the primitive value’s type. For example, putting an int value or variable into object context causes the creation of an Integer object with the value of the int primitive. This coercion is called boxing.

Whereas C++ classes can be defined to have no parent, that is not possible in Java. All Java classes must be subclasses of the root class, Object, or some class that is a descendant of Object. One advantage of this is that some commonly needed methods, such as toString and equals, can be defined in Object and inherited and used by all other classes.

All Java objects are explicit heap dynamic. Most are allocated with the new operator, but there is no explicit deallocation operator. Garbage collection is used for storage reclamation. Like many other language features, although garbage collection avoids some serious problems, such as dangling pointers, it can cause other problems. One such difficulty arises because the garbage
12.4 Support for Object-Oriented Programming in Specific Languages

Collector deallocates, or reclaims the storage occupied by an object, but it does no more. For example, if an object has access to some resource other than heap memory, such as a file or a lock on a shared resource, the garbage collector does not reclaim these. For these situations, Java allows the inclusion of a special method, `finalize`, which is related to a C++ destructor function.

A `finalize` method is implicitly called when the garbage collector is about to reclaim the storage occupied by the object. The problem with `finalize` is that the time it (and the garbage collector) will run cannot be forced or even predicted. The alternative to using `finalize` to reclaim resources held by an object about to be garbage collected is to include a method that does the reclamation. The only problem with this is that all clients of the objects must be aware of this method and remember to call it.

12.4.4.2 Inheritance

In Java, a method can be defined to be `final`, which means that it cannot be overridden in any descendant class. When the `final` reserved word is specified on a class definition, it means the class cannot be subclassed. All of the methods in a final class are implicitly final, which means that the bindings of method calls to the methods of the class are statically bound.

The advantage of defining a class to be final is that no changes to the class are allowed. For example, `String` is a final class and because of that any method that receives a `String` reference in a parameter can depend on the stability of the meaning of `String`’s methods. The disadvantage is that defining a class to be final disallows reuses that require even minor modifications.

Java includes the annotation `@Override`, which informs the compiler to check to determine whether the following method overrides a method in an ancestor class. If it does not, the compiler issues an error message.

Like C++, Java requires that parent class constructor be called before the subclass constructor is called. If parameters are to be passed to the parent class constructor, that constructor must be explicitly called, as in the following example:

```java
super(100, true);
```

If there is no explicit call to the parent class constructor, the compiler inserts a call to the zero-parameter constructor in the parent class.

Java does not support the private derivations of C++. One can surmise that the Java designers believed that subclasses should be subtypes, which they are not when private derivations are supported. Thus, they did not include them. So, Java’s subclasses can be subtypes.

Early versions of Java included a collection, `Vector`, which included a long list of methods for manipulating data in a collection construct. These versions of Java also included a subclass of `Vector`, `Stack`, which added methods for push and pop operations. Unfortunately, because Java does not have private derivation, all of the methods of `Vector` were also visible in the `Stack` class,
which made Stack objects liable to a variety of operations that could invalidate those objects.

Java directly supports only single inheritance. However, it includes a kind of abstract class, called an interface, which provides partial support for multiple inheritance. An interface definition is similar to a class definition, except that it can contain only named constants and method declarations (not definitions). It cannot contain constructors, nonabstract methods, or variable declarations. So, an interface is no more than what its name indicates—it defines only the specification of a class. (Recall that a C++ abstract class can have instance variables and all but one of the methods can be completely defined.) A class does not inherit an interface; it implements it. In fact, a class can implement any number of interfaces. To implement an interface, the class must implement all of the methods whose specifications (but not bodies) appear in the interface definition.

An interface can be used to simulate multiple inheritance. A class can be derived from a class and implement an interface, with the interface taking the place of a second parent class. This is sometimes called mixin inheritance, because the constants and methods of the interface are mixed in with the methods and data inherited from the superclass, as well as any new data and/or methods defined in the subclass.

One more interesting capability of interfaces is that they provide another kind of polymorphism. This is because interfaces can be treated as types. For example, a method can specify a formal parameter that is an interface. Such a formal parameter can accept an actual parameter of any class that implements the interface, making the method polymorphic.

A nonparameter variable also can be declared to be of the type of an interface. Such a variable can reference any object of any class that implements the interface.

One of the problems with multiple inheritance occurs when a class is derived from two parent classes and both define a public method with the same name and protocol. This problem is avoided with interfaces. Although a class that implements an interface must provide definitions for all of the methods specified in the interface, if the class and the interface both include methods with the same name and protocol, the class need not reimplement that method. So, the method name conflicts that can occur with multiple inheritance cannot occur with single inheritance and interfaces. Furthermore, variable name conflicts are completely avoided because interfaces cannot define variables.

An interface is not a replacement for multiple inheritance, because in multiple inheritance there is code reuse, while interfaces provide no code reuse. This is an important difference, because code reuse is one of the primary benefits of inheritance. Java provides one way to partially avoid this deficiency. One of the implemented interfaces could be replaced by an abstract class, which could include code that could be inherited, thereby providing some code reuse.

One problem with interfaces being a replacement for multiple inheritance is the following: If a class attempts to implement two interfaces and both define
methods that have the same name and protocol, there is no way to implement both in the class.

As an example of an interface, consider the `sort` method of the standard Java class, `Array`. Any class that uses this method must provide an implementation of a method to compare the elements to be sorted. The generic `Comparable` interface provides the protocol for this comparing method, which is named `compareTo`. The code for the `Comparable` interface is as follows:

```java
public interface Comparable <T> {
    public int compareTo(T b);
}
```

The `compareTo` method must return a negative integer if the object through which it is called belongs before the parameter object, zero if they are equal, and a positive integer if the parameter belongs before the object through which `compareTo` was called. A class that implements the `Comparable` interface can sort the contents of any array of objects of the generic type, as long as the implemented `compareTo` method for the generic type is implemented and provides the appropriate value.

In addition to interfaces, Java also supports abstract classes, similar to those of C++. The abstract methods of a Java abstract class are represented as just the method's header, which includes the `abstract` reserved word. The abstract class is also marked `abstract`. Of course, abstract classes cannot be instantiated.

Chapter 14 illustrates the use of interfaces in Java event handling.

### 12.4.4.3 Dynamic Binding

In C++, a method must be defined as virtual to allow dynamic binding. In Java, all method calls are dynamically bound unless the called method has been defined as `final`, in which case it cannot be overridden and all bindings are static. Static binding is also used if the method is `static` or `private`, both of which disallow overriding.

### 12.4.4.4 Nested Classes

Java has several varieties of nested classes, all of which have the advantage of being hidden from all classes in their package, except for the nesting class. Non-static classes that are nested directly in another class are called `inner classes`. Each instance of an inner class must have an implicit pointer to the instance of its nesting class to which it belongs. This gives the methods of the nested class access to all of the members of the nesting class, including the private members. Static nested classes do not have this pointer, so they cannot access members of the nesting class. Therefore, static nested classes in Java are like the nested classes of C++.

Though it seems odd in a static-scoped language, the members of the inner class, even the private members, are accessible in the outer class. Such
references must include the variable that references the inner class object. For example, suppose the outer class creates an instance of the inner class with the following statement:

```java
myInner = this.new Inner();
```

Then, if the inner class defines a variable named `sum`, it can be referenced in the outer class as `myInner.sum`.

An instance of a nested class can only exist within an instance of its nesting class. Nested classes can also be anonymous. Anonymous nested classes have complex syntax but are really only an abbreviated way to define a class that is used from just one location. An example of an anonymous nested class appears in Chapter 14.

A local nested class is defined in a method of its nesting class. Local nested classes are never defined with an access specifier (private or public). Their scope is always limited to their nesting class. A method in a local nested class can access the variables defined in its nesting class and the final variables defined in the method in which the local nested class is defined. The members of a local nested class are visible only in the method in which the local nested class is defined.

### 12.4.4.5 Evaluation

Java’s design for supporting object-oriented programming is similar to that of C++, but it employs more consistent adherence to object-oriented principles. Java does not allow parentless classes and uses dynamic binding as the “normal” way to bind method calls to method definitions. This, of course, increases execution time slightly over languages in which many method bindings are static. At the time this design decision was made, however, most Java programs were interpreted, so interpretation time made the extra binding time insignificant. Access controls for the contents of a class definition are rather simple when compared with the jungle of access controls of C++, ranging from derivation controls to friend functions. Finally, Java uses interfaces to provide a form of support for multiple inheritance, which does not have all of the drawbacks of actual multiple inheritance.

### 12.4.5 C#

C#’s support for object-oriented programming is similar to that of Java.

### 12.4.5.1 General Characteristics

C# includes both classes and structs, with the classes being very similar to Java’s classes and the structs being somewhat less powerful constructs. One important difference is that structs are value types; that is, they are stack dynamic. This could cause the problem of object slicing, but this is prevented by the restriction that structs cannot be subclassed. More details of how C# structs differ from its classes appeared in Chapter 11.
12.4.5.2 Inheritance

C# uses the syntax of C++ for defining classes. For example,

```csharp
public class NewClass : ParentClass { . . . }
```

A method inherited from the parent class can be replaced in the derived class by marking its definition in the subclass with `new`. The `new` method hides the method of the same name in the parent class to normal access. However, the parent class version can still be called by prefixing the call with `base`. For example,

```csharp
base.Draw();
```

As with Java, subclasses can be subtypes. C#’s support for interfaces is the same as that of Java. It does not support multiple inheritance.

12.4.5.3 Dynamic Binding

To allow dynamic binding of method calls to methods in C#, both the base method and its corresponding methods in derived classes must be specially marked. The base class method must be marked with `virtual`, as in C++. To make clear the intent of a method in a subclass that has the same name and protocol as a virtual method in an ancestor class, C# requires that such methods be marked `override` if they are to override the parent class virtual method. For example, the C# version of the C++ Shape class that appears in Section 12.4.2.3 is as follows:

```csharp
public class Shape {
    public virtual void Draw() { . . . }
    . . .
}
public class Circle : Shape {
    public override void Draw() { . . . }
    . . .
}
public class Rectangle : Shape {
    public override void Draw() { . . . }
    . . .
}
public class Square : Rectangle {
    public override void Draw() { . . . }
    . . .
}
```

C# includes abstract methods similar to those of C++, except that they are specified with different syntax. For example, the following is a C# abstract method:

```csharp
abstract public void Draw();
```

---

14. Recall that this can be specified in Java with the annotation `@Override`.
A class that includes at least one abstract method is an abstract class, and every abstract class must be marked `abstract`. Abstract classes cannot be instantiated. It follows that any subclass of an abstract class that will be instantiated must implement all abstract methods that it inherits.

As with Java, all C# classes are ultimately derived from a single root class, `Object`. The `Object` class defines a collection of methods, including `ToString`, `Finalize`, and `Equals`, which are inherited by all C# types.

### 12.4.5.4 Nested Classes

A C# class that is directly nested in a class behaves like a Java static nested class (which is like a nested class in C++). Like C++, C# does not support nested classes that behave like the nonstatic nested classes of Java.

### 12.4.5.5 Evaluation

Because C# is a recently designed C-based object-oriented language, one should expect that its designers learned from their predecessors and duplicated the successes of the past and remedied some of the problems. One result of this, coupled with the few problems with Java, is that the differences between C#'s support for object-oriented programming and that of Java are relatively minor. The availability of structs in C#, which Java does not have, can be considered an improvement. Like that of Java, C#'s support for object-oriented programming is simpler than that of C++, which many consider an improvement.

### 12.4.6 Ruby

As stated previously, Ruby is a pure object-oriented programming language in the sense of Smalltalk. Virtually everything in the language is an object and all computation is accomplished through message passing. Although programs have expressions that use infix operators and therefore have the same appearance as expressions in languages like Java, those expressions actually are evaluated through message passing. As is the case with Smalltalk, when one writes `a + b`, it is evaluated by sending the message `+` to the object referenced by `a`, passing a reference to the object `b` as a parameter. In other words, `a + b` is implemented as `a. + b`.

### 12.4.6.1 General Characteristics

Ruby class definitions differ from those of languages such as C++ and Java in that they are executable. Because of this, they are allowed to remain open during execution. A program can add members to a class any number of times, simply by providing secondary definitions of the class that include the new members. During execution, the current definition of a class is the union of all definitions of the class that have been executed. Method definitions are
also executable, which allows a program to choose between two versions of a method definition during execution, simply by putting the two definitions in the then and else clause of a selection construct.

Ruby objects are created with new, which implicitly calls a constructor. The usual constructor in a Ruby class is named initialize. A constructor in a subclass can initialize the data members of the parent class that have setters defined. This is done by calling super with the initial values as actual parameters. super calls the method in the parent class that has the same name as the method in which the call to super appears.

Ruby classes can be nested, but the nested class has no special access to the variables or methods of the nesting class.

All variables in Ruby are references to objects, and all are typeless. Recall that the names of all instance variables in Ruby begin with an at sign (@).

In a clear departure from the other common programming languages, access control in Ruby is different for data than it is for methods. All instance data has private access by default, and that cannot be changed. Therefore, no subclass in Ruby is a subtype. If external access to an instance variable is required, accessor methods must be defined. For example, consider the following skeletal class definition:

```ruby
class MyClass
    # A constructor
    def initialize
        @one = 1
        @two = 2
    end

    # A getter for @one
    def one
        @one
    end

    # A setter for @one
    def one=(my_one)
        @one = my_one
    end

end  # of class MyClass
```

The equal sign (=) attached to the name of the setter method means that its variable is assignable. So, all setter methods have equal signs attached to their names. The body of the one getter method illustrates the Ruby design of methods returning the value of the last expression evaluated when there is no return statement. In this case, the value of @one is returned.

Because getter and setter methods are so frequently needed, Ruby provides shortcuts for creating them. If one wants a class to have getter methods for the
two instance variables, `@one` and `@two`, those getters can be specified with the single statement in the class:

```
attr_reader :one, :two
```

`attr_reader` is actually a function call, using `:one` and `:two` as the actual parameters. Preceding a variable with a colon (:) causes the variable name to be used, rather than dereferencing it to the object to which it refers. Instead of passing a value or an address, the text of the variable's name is passed. This is exactly how macro parameters are passed.

The function that similarly creates setters is called `attr_writer`. This function has the same parameter profile as `attr_reader`.

The functions for creating getter and setter methods are so named because they provide the protocol for objects of the class, which then are called attributes. So, the attributes of a class define the data interface (the data made public through accessor methods) to objects of the class.

Class variables, which are specified by preceding their names with two at signs (`@@`), are private to the class and its instances. That privacy cannot be changed. Also, unlike global and instance variables, class variables must be initialized before they are used.

### 12.4.6.2 Inheritance

Subclasses are defined in Ruby using the less-than symbol (<), rather than the colon of C++. For example,

```
class MySubClass < BaseClass
```

One distinct thing about the method access controls of Ruby is that they can be changed in a subclass, simply by calling the access control functions. This means that two subclasses of a base class can be defined so that objects of one of the subclasses can access a method defined in the base class, but objects of the other subclass cannot. Also, this allows one to change the access of a publicly accessible method in the base class to a privately accessible method in the subclass.

### 12.4.6.3 Dynamic Binding

Support for dynamic binding in Ruby is the same as it is in Smalltalk. Variables are not typed; rather, they are all references to objects of any class. So, all variables are polymorphic and all bindings of method calls to methods are dynamic.

### 12.4.6.4 Evaluation

Because Ruby is an object-oriented programming language in the purest sense, its support for object-oriented programming is obviously adequate. However, access control to class members is weaker than that of C++. Ruby does not
### Table 12.1 Designs

<table>
<thead>
<tr>
<th>DESIGN ISSUE/LANGUAGE</th>
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<th>OBJECTIVE-C</th>
<th>JAVA</th>
<th>C#</th>
<th>RUBY</th>
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</thead>
<tbody>
<tr>
<td>Exclusivity of objects</td>
<td>All data are objects</td>
<td>Primitive types plus objects</td>
<td>Primitive types plus objects</td>
<td>Primitive types plus objects</td>
<td>Primitive types plus objects</td>
<td>All data are objects</td>
</tr>
<tr>
<td>Are subclasses subtypes?</td>
<td>They can be and usually are</td>
<td>They can be and usually are</td>
<td>They can be and usually are</td>
<td>They can be and usually are</td>
<td>They can be and usually are</td>
<td>No subclasses are subtypes</td>
</tr>
<tr>
<td>Single and multiple inheritance</td>
<td>Single only</td>
<td>Both</td>
<td>Single only, but some effects with protocols</td>
<td>Single only, but some effects with interfaces</td>
<td>Single only, but some effects with interfaces</td>
<td>Single only, but some effects with modules</td>
</tr>
<tr>
<td>Allocation and deallocation of objects</td>
<td>All objects are heap allocated; allocation is explicit and deallocation is implicit</td>
<td>Objects can be static, stack dynamic, or heap dynamic; allocation and deallocation are explicit</td>
<td>All objects are heap dynamic; allocation is explicit and deallocation is implicit</td>
<td>All objects are heap dynamic; allocation is explicit and deallocation is implicit</td>
<td>All objects are heap dynamic; allocation is explicit and deallocation is implicit</td>
<td>All objects are heap dynamic; allocation is explicit and deallocation is implicit</td>
</tr>
<tr>
<td>Dynamic and static binding</td>
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<td>Method binding can be either</td>
<td>Method binding can be either</td>
<td>Method binding can be either</td>
<td>Method binding can be either</td>
<td>All method bindings are dynamic</td>
</tr>
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<td>Nested classes?</td>
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<td>Yes</td>
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<td>Yes</td>
</tr>
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<td>Initialization</td>
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<td>Constructors are implicitly called</td>
<td>Constructors are implicitly called</td>
<td>Constructors are implicitly called</td>
</tr>
</tbody>
</table>
support abstract classes or interfaces, although its mixins are closely related to interfaces. Finally, in large part because Ruby is interpreted, its execution efficiency is far worse than that of the compiled languages.

Table 12.1 summarizes how the designers of the languages in this section chose to deal with the design issues described in Section 12.3

12.5 Implementation of Object-Oriented Constructs

There are at least two parts of language support for object-oriented programming that pose interesting questions for language implementers: storage structures for instance variables and the dynamic bindings of messages to methods. In this section, we provide a brief look at these.

12.5.1 Instance Data Storage

In C++, classes are defined as extensions of C’s record structures—structs. This similarity suggests a storage structure for the instance variables of class instances—that of a record. This form of this structure is called a class instance record (CIR). The structure of a CIR is static, so it is built at compile time and used as a template for the creation of the data of class instances. Every class has its own CIR. When a derivation takes place, the CIR for the subclass is a copy of that of the parent class, with entries for the new instance variables added at the end.

Because the structure of the CIR is static, access to all instance variables can be done as it is in records, using constant offsets from the beginning of the CIR instance. This makes these accesses as efficient as those for the fields of records.

12.5.2 Dynamic Binding of Method Calls to Methods

Methods in a class that are statically bound need not be involved in the CIR for the class. However, methods that will be dynamically bound must have entries in this structure. Such entries could simply have a pointer to the code of the method, which must be set at object creation time. Calls to a method could then be connected to the corresponding code through this pointer in the CIR. The drawback to this technique is that every instance would need to store pointers to all dynamically bound methods that could be called from the instance.

Notice that the list of dynamically bound methods that can be called from an instance of a class is the same for all instances of that class. Therefore, the list of such methods must be stored only once. So the CIR for an instance needs only a single pointer to that list to enable it to find called methods. The storage structure for the list is often called a virtual method table (vtable). Method calls can be represented as offsets from the beginning of the vtable. Polymorphic variables of an ancestor class always reference the CIR of the correct type object, so getting to the correct version of a dynamically bound method is assured. Consider the following Java example, in which all methods are dynamically bound:
The CIRs for the A and B classes, along with their vtables, are shown in Figure 12.7. Notice that the method pointer for the area method in B’s vtable points to the code for A’s area method. The reason is that B does not override A’s area method, so if a client of B calls area, it is the area method inherited from A. On the other hand, the pointers for draw and sift in B’s vtable point to B’s draw and sift. The draw method is overridden in B and sift is defined as an addition in B.

Multiple inheritance complicates the implementation of dynamic binding. Consider the following three C++ class definitions:

class A {
    public:
        int a;
        virtual void fun() { . . . }
        virtual void init() { . . . }
};

class B extends A {
    public int c, d;
    public void draw() { . . . }
    public void sift() { . . . }
}

![Figure 12.7](image)

An example of the CIRs with single inheritance
The C class inherits the variable a and the init method from the A class. It redefines the fun method, although both its fun and that of the parent class A are potentially visible through a polymorphic variable (of type A). From B, C inherits the variable b and the sum method. C defines its own variable, c, and defines an uninherited method, dud. A CIR for C must include A's data, B's data, and C's data, as well as some means of accessing all visible methods. Under single inheritance, the CIR would include a pointer to a vtable that has the addresses of the code of all visible methods. With multiple inheritance, however, it is not that simple. There must be at least two different views available in the CIR—one for each of the parent classes, one of which includes the view for the subclass, C. This inclusion of the view of the subclass in the parent class's view is just as in the implementation of single inheritance.

There must also be two vtables: one for the A and C view and one for the B view. The first part of the CIR for C in this case can be the C and A view, which begins with a vtable pointer for the methods of C and those inherited from A, and includes the data inherited from A. Following this in C's CIR is the B view part, which begins with a vtable pointer for the virtual methods of B, which is followed by the data inherited from B and the data defined in C. The CIR for C is shown in Figure 12.8.

Figure 12.8
An example of a subclass CIR with multiple parents
12.6 Reflection

A discussion of reflection is not a perfect fit into a chapter on object orientation, but it is even a worse fit into any other chapter of this book. So, this is where we put it.

12.6.1 Introduction

In general, the later bindings take place in a programming language, the more flexible the language is. For example, the late binding of data types in scripting languages and functional languages allows their programs to be more generic than those in the static-typed languages. Likewise, the dynamic binding of method calls to methods that is part of the object-oriented languages allows their programs to be easier to maintain and extend. Among other things, reflection provides the possibility of late binding of calls to methods that are outside the inheritance hierarchy of the calling code.

12.6.2 What Is Reflection?

A programming language that supports reflection allows its programs to have run-time access to their types and structure and to be able to dynamically modify their behavior. To allow a program to examine its types and structure, that information must be gathered by the compiler or interpreter and made available to the program. Just as information about the structure of a database is called metadata, the types and structure of a program are called metadata. The process of a program examining its metadata is called introspection. A program can modify its behavior dynamically in several different ways: it could change its metadata directly, it could use the metadata, or it could intercede in the execution of the program. The first of these is complicated; the second is less complex and is common among languages; the third often is called intercession.

Some of the primary uses of reflection are in the construction of software tools. A class browser needs to enumerate the classes of a program. Visual Integrated Development Environments can use type information to assist a developer in building type-correct code. Debuggers must be able to examine private fields and methods of classes. Test systems need to be able to discover all of the methods of a class to be sure that test data drives all of them.

To illustrate a relatively simple and common use of reflection, we pose the following problem. A zoo has a large area devoted to birds. The flight cage for each species includes a plaque that provides general information about the inhabitant’s species. Included on the plaque is a small screen onto which a visitor with special interest in the species can enter his or her entrance ticket number. At the exit to the bird exhibit, the visitor can again enter his or her ticket number on a small screen, which causes a computer to print pictures of the birds for which the visitor previously indicated particular interest. The computer system that
supports these activities has an object that includes a method that draws a picture of its bird for each of the birds on display. This seems simple enough. When a visitor selects a bird at its flight cage, the system places a reference to an object associated with that bird in a list. At the exit of the exhibit, the system calls the draw method of each object in the visitor’s list. The process is complicated by the fact that the zoo supplies only some of the bird objects. Some of them are purchased from third-party vendors and some are donated by zoo benefactors. Because of the multiple sources of the bird objects, they do not have a common base class (other than Object) and do not implement a common interface, so what type references can be saved? One obvious solution is to make each bird object the subclass of a base class. References of the base class type could be stored in the list and dynamic binding could be used to invoke the draw methods. The drawback of this approach is that every bird class would need to be modified to make the new classes subclasses of the common base class. It would be better if the new bird classes could simply be added to a code file without modification. Another possible solution would be to use instanceof and casting to determine the concrete types of the references. This would add much code to the system, increasing its complexity and cost of maintenance. A better solution is to use the dynamic binding that is possible with reflection.

### 12.6.3 Reflection in Java

Java provides limited support for reflection. The primary class of the metadata is defined in the namespace, `java.lang.Class`. This has the unfortunately confusing name, `Class`. The Java run-time system instantiates an instance of `Class` for each object in the program. The `Class` class provides a collection of methods to examine the type information and members of the program objects. `Class` is the access point for all of the reflection API.

If the program has a reference to an object (not a primitive), the `Class` object of that object can be obtained by calling its `getClass` method. All classes inherit `getClass` from `Object`, from which all objects descend. Consider the following examples:

```java
float[] totals = new float[100];
Class fltlist = totals.getClass();
Class stg = “hello”.getClass();
```

The value of the variable `fltlist` will be the `Class` object of the `totals` array object. The value of `stg` will be the `Class` object of `String` (because “hello” is an instance of `String`).

If there is no object of a class, its `Class` object can be obtained through the class’ name by attaching `.class` to the name. For example, we could have the following:

```java
Class stg = String.class;
```

---

15. It is so named because its instantiations are themselves classes.
If the class has no name, its `Class` object can still be obtained by attaching `.class` to the class definition. For example, consider the following:

```java
Class intmat = int[][] .class;
```

The `.class` modifier can also be attached to primitive types. Although `float.getClass()` is illegal, `float.class` is not.

There are four methods to get the `Class` of a method. The `getMethod` method searches a class to find a specific public method defined in the class or inherited by the class. The `getMethods` method returns an array of all of the public methods defined in a class or inherited by the class. The `getDeclaredMethod` method searches for a specific method declared in a class, including private methods. The `getDeclaredMethods` method returns all of the methods defined in a class.

If the `Class` object of an object is known and a particular method defined by the class of the object is found, that method can be called through the `Method` object of the method with the `invoke` method. For example, if the `Method` object named `method` is found with `getMethod`, it can be called with the following:

```java
method.invoke(…);
```

We can now develop a solution in Java for the problem posed in Section 12.6.2. The heart of this application is a class that defines a method that is passed an `Object` reference. The method determines the class of the passed reference, finds a `draw` method of that class, and calls that method. The solution class is tested with a second class, `ReflectTest`, which creates an array of three `Object` references to classes that represent three different birds. Each of these defines a `draw` method that, when called, displays a message indicating that it was called. Then the test calls the class method, passing the elements of the array of references.

The caller method can raise three different exceptions, each of which is handled in the method.

```java
// A project to illustrate dynamic method calling
// using reflection in Java
package reflect;
import java.lang.reflect.*;
// A class to test the Reflect class
// Creates three objects that represent different birds
// and calls a method that dynamically calls the draw
// methods of the three bird classes
public class ReflectTest {
    public static void main(String[] args) {
        Object[] birdList = new Object[3];
        birdList[0] = new Bird1();
        birdList[1] = new Bird2();
        birdList[2] = new Bird3();
        Reflect.callDraw(birdList[2]);
        Reflect.callDraw(birdList[0]);
        Reflect.callDraw(birdList[1]);
    }
}
```
class Reflect {
    public static void callDraw(Object birdObj) {
        Class cls = birdObj.getClass();
        try {
            // Find the draw method of the given class
            Method method = cls.getMethod("draw");
            // Dynamically call the method
            method.invoke(birdObj);
        } // In case the given class does not support draw
        catch (NoSuchMethodException e) {
            throw new IllegalArgumentException (
                cls.getName() + "does not support draw");
        } // In case the callDraw cannot call draw
        catch (IllegalAccessException e) {
            throw new IllegalArgumentException ("Insufficient access permissions to call" + 
                "draw in class " + cls.getName());
        } // In case draw throws an exception
        catch (InvocationTargetException e) {
            throw new RuntimeException(e);
        }
    }
}

class Bird1 {
    public void draw() {
        System.out.println("This is draw from Bird1");
    }
}
class Bird2 {
    public void draw() {
        System.out.println("This is draw from Bird2");
    }
}
class Bird3 {
    public void draw() {
        System.out.println("This is draw from Bird3");
    }
}

The output of this program is as follows:

This is the draw from Bird3
This is the draw from Bird1
This is the draw from Bird2
12.6.4 Reflection in C#

Support for reflection in C# is similar to that of Java, with a few important differences. In C#, as in all .NET languages, the compiler places the intermediate code, written in Common Intermediate Language (CIL), in an assembly, which could include several files. An assembly also contains an assembly version number and the metadata for all classes defined in the assembly, as well as for all external classes it uses.

Instead of the java.lang.Class namespace, System.Type is used in .NET; instead of java.lang.reflect, System.Reflection is used. Rather than the getClass method, getType is used to get the class of an instance. Also, the .NET languages use the typeof operator in place of the .class field used in Java. Following is a C# version of the Java project shown above:

```csharp
using System;
using System.Reflection;
namespace TestReflect
{
    // A project to illustrate dynamic method calling
    // using reflection in C#
    // A class to test the Reflect class
    // Creates three objects that represent different birds
    // and calls a method that dynamically calls the draw
    // methods of the three bird classes
    public class ReflectTest
    {
        public static void Main(String[] args)
        {
            Object[] birdList = new Object[3];
birdList[0] = new Bird1();
birdList[1] = new Bird2();
birdList[2] = new Bird3();
            Reflect.callDraw(birdList[2]);
            Reflect.callDraw(birdList[0]);
            Reflect.callDraw(birdList[1]);
        }
    }

    // A class to define the method that dynamically calls the
    // methods of a passed class object
    class Reflect
    {
        public static void callDraw(Object birdObj)
        {
            Type typ = birdObj.GetType();
            // Find the draw method of the given class
            MethodInfo method = typ.GetMethod("draw");
            // Dynamically call the method
            method.Invoke(birdObj, null);
        }
    }

    class Bird1
    {
        public void draw()
        {
```
Console.WriteLine("This is draw from Bird1");
}
}

class Bird2 {
    public void draw() {
        Console.WriteLine("This is draw from Bird2");
    }
}

class Bird3 {
    public void draw() {
        Console.WriteLine("This is draw from Bird3");
    }
}

Our simple example of dynamic method binding shows just one of the many uses of reflection.

In addition to the methods and fields of a class, the following program elements can be accessed with reflection in both Java and C#: class modifiers, such as public, static, and final, constructors, method parameter types, and implemented interfaces. Also, a description of the inheritance path of a class can be introspected. In C#, but not Java, the names of the formal parameters of methods can be discovered.

One significant difference between Java’s reflection and that of C# is the System.Reflection.Emit namespace, which is part of .NET. This namespace provides the ability to create CIL code and an assembly to house that code. Java provides no such capability, although it can be done with tools from other suppliers.

Although reflection adds a variety of capabilities to the static-typed languages Java and C#, the user of reflection must be aware of its downsides:

- Performance nearly always suffers with the use of reflection. Resolving types, methods, and fields at run time are not part of the cost of running nonreflective code. Also, when types are dynamically resolved, some optimizations cannot be done on the code.
- Reflection exposes private fields and methods, which violate the rules of abstraction and information hiding, and also may result in unexpected side effects and adversely affect portability.
- Although the advantage of early type checking is widely accepted, the late binding that is possible with reflection obviously negates that advantage.
- Some reflective operations may not work when the code is run under a security manager, also making it non-portable. One such security environment is that of running applets. In most cases, if a problem can be solved without reflection, reflection should not be used.

Reflection is an integral part of most dynamically typed languages. In LISP, reflection is routinely used and the dynamic construction and execution of code is not uncommon. In other interpreted languages, such as JavaScript, Perl, and
Python, the symbol table is kept during interpretation, providing all useful type information.

In Python, for example, the type method returns the type of a given value. For example, `type([7, 14, 21])` is list. The isinstance method returns a Boolean value if its first parameter has the type named in its second parameter. For example, `isinstance(17, int)` returns True. The callable function is used to determine whether an expression returns a function object. The `dir` function returns the list of attributes, both data and methods, of its parameter object.

**Summary**

Object-oriented programming is based on three fundamental concepts: abstract data types, inheritance, and dynamic binding. Object-oriented programming languages support the paradigm with classes, methods, objects, and message passing.

The discussion of object-oriented programming languages in this chapter revolves around seven design issues: exclusivity of objects, subclasses and subtypes, type checking and polymorphism, single and multiple inheritance, dynamic binding, explicit or implicit deallocation of objects, and nested classes.

Smalltalk is a pure object-oriented language—everything is an object and all computation is accomplished through message passing. All type checking and binding of messages to methods is dynamic, and all inheritance is single. Smalltalk has no explicit deallocation operation.

C++ provides support for data abstraction, inheritance, and optional dynamic binding of messages to methods, along with all of the conventional features of C. This means that it has two distinct type systems. C++ provides multiple inheritance and explicit object deallocation. It includes a variety of access controls for the entities in classes, some of which prevent subclasses from being subtypes. Both constructor and destructor methods can be included in classes; both are usually implicitly called.

While Smalltalk’s dynamic type binding provides somewhat more programming flexibility than the hybrid language C++, it is far less efficient.

Objective-C supports both procedural and object-oriented programming. It is less complex and less widely used than C++. Only single inheritance is supported, although it has categories, which allow mixins of additional methods that can be added to a class. It also has protocols, which are similar to Java’s interfaces. A class can adopt any number of protocols. Constructors can have any name, but they must be explicitly called. Polymorphism is supported with the predefined type, `id`. A variable of `id` type can reference any object. When a method is called through an object referenced by a variable of type `id`, the binding is dynamic.

Unlike C++, Java is not a hybrid language; it is meant to support only object-oriented programming. Java has both primitive scalar types and classes. All objects are allocated from the heap and are accessed through reference variables. There is no explicit object deallocation operation—garbage collection is used. The only subprograms are methods, and they can be called only through
objects or classes. Only single inheritance is directly supported, although a kind of multiple inheritance is possible using interfaces. All binding of messages to methods is dynamic, except in the case of methods that cannot be overridden. In addition to classes, Java includes packages as a second encapsulation construct.

C#, which is based on C++ and Java, supports object-oriented programming. Objects can be instantiated from either classes or structs. The struct objects are stack dynamic and do not support inheritance. Methods in a derived class can call the hidden methods of the parent class by including `base` on the method name. Methods that can be overridden must be marked `virtual`, and the overriding methods must be marked with `override`. All classes (and all primitives) are derived from `Object`.

Ruby is an object-oriented scripting language in which all data are objects. As with Smalltalk, all objects are heap allocated and all variables are typeless references to objects. All constructors are named `initialize`. All instance data are private, but getter and setter methods can be easily included. The collection of all instance variables for which access methods have been provided forms the public interface to the class. Such instance data are called attributes. Ruby classes are dynamic in the sense that they are executable and can be changed at any time. Ruby supports only single inheritance.

The instance variables of a class are stored in a CIR, the structure of which is static. Subclasses have their own CIRs, as well as the CIR of their parent class. Dynamic binding is supported with a virtual method table, which stores pointers to specific methods. Multiple inheritance greatly complicates the implementation of CIRs and virtual method tables.

### REVIEW QUESTIONS

1. Describe the three characteristic features of object-oriented languages.
2. What is the difference between a class variable and an instance variable?
3. What is multiple inheritance?
4. What is a polymorphic variable?
5. What is an overriding method?
6. Describe a situation where dynamic binding is a great advantage over static binding.
7. What is a virtual method?
8. What is an abstract method? What is an abstract class?
9. Describe briefly the seven design issues used in this chapter for object-oriented languages.
10. What is a nesting class?
11. What is the message protocol of an object?
12. From where are Smalltalk objects allocated?
13. Explain how Smalltalk messages are bound to methods. When does this take place?
14. What type checking is done in Smalltalk? When does it take place?
15. What kind of inheritance, single or multiple, does Smalltalk support?
16. What are the two most important effects that Smalltalk has had on computing?
17. In essence, all Smalltalk variables are of a single type. What is that type?
18. From where can C++ objects be allocated?
19. How are C++ heap-allocated objects deallocated?
20. Are all C++ subclasses subtypes? If so, explain. If not, why not?
21. Under what circumstances is a C++ method call statically bound to a method?
22. What drawback is there to allowing designers to specify which methods can be statically bound?
23. What are the differences between private and public derivations in C++?
24. What is a friend function in C++?
25. What is a pure virtual function in C++?
26. How are parameters sent to a superclass’s constructor in C++?
27. What is the single most important practical difference between Smalltalk and C++?
28. If an Objective-C method returns nothing, what return type is indicated in its header?
29. Does Objective-C support multiple inheritance?
30. Can an Objective-C class not specify a parent class in its header?
31. What is the root class in Objective-C?
32. In Objective-C, how can a method indicate that it cannot be overridden in descendant classes?
33. What is the purpose of an Objective-C category?
34. What is the purpose of an Objective-C protocol?
35. What is the primary use of the id type in Objective-C?
36. How is the type system of Java different from that of C++?
37. From where can Java objects be allocated?
38. What is boxing?
39. How are Java objects deallocated?
40. Are all Java subclasses subtypes?
41. How are superclass constructors called in Java?
42. Under what circumstances is a Java method call statically bound to a method?
43. In what way do overriding methods in C# syntactically differ from their counterparts in C++?
44. How can the parent version of an inherited method that is overridden in a subclass be called in that subclass in C#?
45. How does Ruby implement primitive types, such as those for integer and floating-point data?
46. How are getter methods defined in a Ruby class?
47. What access controls does Ruby support for instance variables?
48. What access controls does Ruby support for methods?
49. Are all Ruby subclasses subtypes?
50. Does Ruby support multiple inheritance?
51. What does reflection allow a program to do?
52. In the context of reflection, what is metadata?
53. What is introspection?
54. What is intercession?
55. What class in Java stores information about classes in a program?
56. For what is the Java name extension .class used?
57. What does the Java getMethods method do?
58. For what is the C# namespace System.Reflection.Emit used?

**Problem Set**

1. What important part of support for object-oriented programming is missing in Java?
2. Explain the principle of abstraction.
3. Explain the problem associated with the reuse of abstract data types.
5. Compare the class entity access controls of C++ and Java.
6. Compare the multiple inheritance of C++ with that provided by interfaces in Java.
7. What advantages does a derived class have over its base class?
8. Explain the two problems with object-oriented programming that are ameliorated by structure-oriented programming.
9. Describe the categories of changes that a subclass can make to its parent class.
10. Explain one advantage of inheritance.
11. Explain the advantages and disadvantages of having the friend function in object-oriented programming languages.
12. What exactly does it mean for a subclass to have an is-a relationship with its parent class?

13. Describe the issue of how closely the parameters of an overloading method must match those of the method it overloads.

14. Explain diamond inheritance.

15. The designers of Java obviously thought it was not worth the additional efficiency of allowing any method to be statically bound, as is the case with C++. What are the arguments for and against the Java design?

16. What is the primary reason why all Java objects have a common ancestor?

17. What is the purpose of the new operator in C++?

18. What would be gained if Java allowed stack-dynamic objects as well as heap-dynamic objects? What would be the disadvantage of having both?

19. What are the differences between a C++ abstract class and a Java interface?

20. Compare the support for different programming paradigms in C++ with that of Smalltalk.

21. Compare the capabilities and use of Objective-C protocols with Java’s interfaces.

22. Critically evaluate the decision by the designers of Objective-C to use Smalltalk’s syntax for method calls, rather than the conventional syntax used by most imperative-based languages that support object-oriented programming.

23. Explain why allowing a class to implement multiple interfaces in Java and C# does not create the same problems that multiple inheritance in C++ creates.

24. Study and explain the issue of why Java does not include C++’s destructor even though both these languages have constructors.

25. What mechanism should be used to define an abstract class in C++?

26. Compare the access mechanisms of standalone functions and friend functions.

27. Compare the type error detection for instance variables in Java and Ruby.

28. Explain the concept of object slicing.

---

**Programming Exercises**

1. Rewrite the single_linked_list, stack_2, and queue_2 classes in Section 12.5.2 in Java and compare the result with the C++ version in terms of readability and ease of programming.

2. Repeat Programming Exercise 1 using Ruby.

3. Repeat Programming Exercise 1 using Objective-C.
4. Design and implement a C++ program that defines a base class A, which has a subclass B, which itself has a subclass C. The A class must implement a method, which is overridden in both B and C. You must also write a test class that instantiates A, B, and C and includes three calls to the method. One of the calls must be statically bound to A’s method. One call must be dynamically bound to B’s method, and one must be dynamically bound to C’s method. All of the method calls must be through a pointer to class A.

5. Write a program in C++ that calls both a dynamically bound method and a statically bound method a large number of times, timing the calls to both of the two. Compare the timing results and compute the difference of the time required by the two. Explain the results.

Concurrency

13.1 Introduction
13.2 Introduction to Subprogram-Level Concurrency
13.3 Semaphores
13.4 Monitors
13.5 Message Passing
13.6 Ada Support for Concurrency
13.7 Java Threads
13.8 C# Threads
13.9 Concurrency in Functional Languages
13.10 Statement-Level Concurrency
Chapter 13  Concurrency

This chapter begins with introductions to the various kinds of concurrency at the subprogram, or unit level, and at the statement level. Included is a brief description of the most common kinds of multiprocessor computer architectures. Next, a lengthy discussion on unit-level concurrency is presented. This begins with a description of the fundamental concepts that must be understood before discussing the problems and challenges of language support for unit-level concurrency, specifically competition and cooperation synchronization. Next, the design issues for providing language support for concurrency are described. Following this is a detailed discussion of three major approaches to language support for concurrency: semaphores, monitors, and message passing. A pseudocode example program is used to demonstrate how semaphores can be used. Ada and Java are used to illustrate monitors; for message passing, Ada is used. The Ada features that support concurrency are described in some detail. Although tasks are the focus, protected objects (which are effectively monitors) are also discussed. Support for unit-level concurrency using threads in Java and C# is then discussed, including approaches to synchronization. This is followed by brief overviews of support for concurrency in several functional programming languages. The last section of the chapter is a brief discussion of statement-level concurrency, including an introduction to part of the language support provided for it in High-Performance Fortran.

13.1 Introduction

Concurrency in software execution can occur at four different levels: instruction level (executing two or more machine instructions simultaneously), statement level (executing two or more high-level language statements simultaneously), unit level (executing two or more subprogram units simultaneously), and program level (executing two or more programs simultaneously). Because no language design issues are involved with them, instruction-level and program-level concurrency are not discussed in this chapter. Concurrency at both the subprogram and the statement levels is discussed, with most of the focus on the subprogram level.

At first glance, concurrency may appear to be a simple concept, but it presents significant challenges to the programmer, the programming language designer, and the operating system designer (because much of the support for concurrency is provided by the operating system).

Concurrent control mechanisms increase programming flexibility. They were originally invented to be used for particular problems faced in operating systems, but they are required for a variety of other programming applications. One of the most commonly used programs is Web browsers, whose design is based heavily on concurrency. Browsers must perform many different functions at the same time, among them sending and receiving data from Web servers, rendering text and images on the screen, and reacting to user actions with the mouse and the keyboard. Some contemporary browsers, for example Internet Explorer 9, use the extra core processors that are part of many contemporary personal computers to perform some of their processing, for example the
interpretation of client-side scripting code. Another example is the software systems that are designed to simulate actual physical systems that consist of multiple concurrent subsystems. For all of these kinds of applications, the programming language (or a library or at least the operating system) must support unit-level concurrency.

Statement-level concurrency is quite different from concurrency at the unit level. From a language designer’s point of view, statement-level concurrency is largely a matter of specifying how data should be distributed over multiple memories and which statements can be executed concurrently.

The goal of developing concurrent software is to produce scalable and portable concurrent algorithms. A concurrent algorithm is scalable if the speed of its execution increases when more processors are available. This is important because the number of processors sometimes increases with the new generations of machines. The algorithms must be portable because the lifetime of hardware is relatively short. Therefore, software systems should not depend on a particular architecture—that is, they should run efficiently on machines with different architectures.

The intention of this chapter is to discuss the aspects of concurrency that are most relevant to language design issues, rather than to present a definitive study of all of the issues of concurrency, including the development of concurrent programs. That would clearly be inappropriate for a book on programming languages.

### 13.1.1 Multiprocessor Architectures

A large number of different computer architectures have more than one processor and can support some form of concurrent execution. Before beginning to discuss concurrent execution of programs and statements, we briefly describe some of these architectures.

The first computers that had multiple processors had one general-purpose processor and one or more other processors, often called peripheral processors, that were used only for input and output operations. This architecture allowed those computers, which appeared in the late 1950s, to execute one program while concurrently performing input or output for other programs.

By the early 1960s, there were machines that had multiple complete processors. These processors were used by the job scheduler of the operating system, which distributed separate jobs from a batch-job queue to the separate processors. Systems with this structure supported program-level concurrency.

Machines that appeared in the mid-1960s had several identical partial processors. These were fed instructions from a single instruction stream. For example, some machines had two or more floating-point multipliers, while others had two or more complete floating-point arithmetic units. The compilers for these machines were required to determine which instructions could be executed concurrently and to schedule these instructions accordingly. Systems with this structure supported instruction-level concurrency.
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In 1966, Michael J. Flynn suggested a categorization of computer architectures defined by whether the instruction and data streams were single or multiple. The names of these were widely used from the 1970s to the early 2000s. The two categories that used multiple data streams are defined as follows: Computers that have multiple processors that execute the same instruction simultaneously, each on different data, are called Single Instruction, Multiple Data (SIMD) architecture computers. In an SIMD computer, each processor has its own local memory. One processor controls the operation of the other processors. Because all of the processors, except the controller, execute the same instruction at the same time, no synchronization is required in the software. Perhaps the most widely used SIMD machines are a category of machines called vector processors. They have groups of registers that store the operands of a vector operation in which the same instruction is executed on the whole group of operands simultaneously. Originally, the kinds of programs that could most benefit from this architecture were in scientific computation, an area of computing that is often the target of multiprocessor machines. However, SIMD processors are now used for a variety of application areas, among them graphics and video processing. Until recently, most supercomputers were vector processors.

Computers that have multiple processors that operate independently but whose operations can be synchronized are called Multiple Instruction, Multiple Data (MIMD) computers. Each processor in an MIMD computer executes its own instruction stream. MIMD computers can appear in two distinct configurations: distributed and shared memory systems. The distributed MIMD machines, in which each processor has its own memory, can be either built in a single chassis or distributed, perhaps over a large area. The shared-memory MIMD machines obviously must provide some means of synchronization to prevent memory access clashes. Even distributed MIMD machines require synchronization to operate together on single programs. MIMD computers, which are more general than SIMD computers, support unit-level concurrency. The primary focus of this chapter is on language design for shared memory MIMD computers, which are often called multiprocessors.

With the advent of powerful but low-cost single-chip computers, it became possible to have large numbers of these microprocessors connected into physically small networks within a single chassis. These kinds of computers, which often use off-the-shelf microprocessors, have appeared from a number of different manufacturers.

One important reason why software has not evolved faster to make use of concurrent machines is that the power of processors has continually increased. One of the strongest motivations to use concurrent machines is to increase the speed of computation. However, two hardware factors have combined to provide faster computation, without requiring any change in the architecture of software systems. First, processor clock rates have become faster with each new generation of processors (the generations have appeared roughly every 18 months). Second, several different kinds of concurrency have been built into the processor architectures. Among these are the pipelining of instructions and data from the memory to the processor (instructions are fetched and decoded
for future execution while the current instruction is being executed), the use of separate lines for instructions and data, prefetching of instructions and data, and parallelism in the execution of arithmetic operations. All of these are collectively called hidden concurrency. The result of the increases in execution speed is that there have been great productivity gains without requiring software developers to produce concurrent software systems.

However, the situation is now changing. The end of the sequence of significant increases in the speed of individual processors is now near. Significant increases in computing power now result from increases in the number of processors, for example large server systems like those run by Google and Amazon and scientific research applications. Many other large computing tasks are now run on machines with large numbers of relatively small processors.

Another recent advance in computing hardware was the development of multiple processors on a single chip, such as with the Intel Core Duo and Core Quad chips, which is putting more pressure on software developers to make more use of the available multiple processor machines. If they do not, the concurrent hardware will be wasted and productivity gains will significantly diminish.

### 13.1.2 Categories of Concurrency

There are two distinct categories of concurrent unit control. The most natural category of concurrency is that in which, assuming that more than one processor is available, several program units from the same program literally execute simultaneously. This is physical concurrency. A slight relaxation of this concept of concurrency allows the programmer and the application software to assume that there are multiple processors providing actual concurrency, when in fact the actual execution of programs is taking place in interleaved fashion on a single processor. This is logical concurrency. From the programmer’s and language designer’s points of view, logical concurrency is the same as physical concurrency. It is the language implementor’s task, using the capabilities of the underlying operating system, to map the logical concurrency to the host hardware. Both logical and physical concurrencies allow the concept of concurrency to be used as a program design methodology. For the remainder of this chapter, the discussion will apply to both physical and logical concurrencies.

One useful technique for visualizing the flow of execution through a program is to imagine a thread laid on the statements of the source text of the program. Every statement reached on a particular execution is covered by the thread representing that execution. Visually following the thread through the source program traces the execution flow through the executable version of the program. Of course, in all but the simplest of programs, the thread follows a highly complex path that would be impossible to follow visually. Formally, a thread of control in a program is the sequence of program points reached as control flows through the program.

Programs that have coroutines (see Chapter 9) but no concurrent subprograms, though they are sometimes called quasi-concurrent, have a single
thread of control. Programs executed with physical concurrency can have multiple threads of control. Each processor can execute one of the threads. Although logically concurrent program execution may actually have only a single thread of control, such programs can be designed and analyzed only by imagining them as having multiple threads of control. A program designed to have more than one thread of control is said to be *multithreaded*. When a multithreaded program executes on a single-processor machine, its threads are mapped onto a single thread. It becomes, in this scenario, a virtually multithreaded program.

Statement-level concurrency is a relatively simple concept. In a common use of statement-level concurrency, loops that include statements that operate on array elements are unwound so that the processing can be distributed over multiple processors. For example, a loop that executes 500 repetitions and includes a statement that operates on one of 500 array elements may be unwound so that each of 10 different processors can simultaneously process 50 of the array elements.

### 13.1.3 Motivations for the Use of Concurrency

There are at least four different reasons to design concurrent software systems. The first reason is the speed of execution of programs on machines with multiple processors. These machines provide an effective way of increasing the execution speed of programs, provided that the programs are designed to make use of the concurrent hardware. There are now a large number of installed multiple-processor computers, including many of the personal computers sold in the last few years. It is wasteful not to use this hardware capability.

The second reason is that even when a machine has just one processor, a program written to use concurrent execution can be faster than the same program written for sequential (nonconcurrent) execution. The requirement for this to happen is that the program is not compute bound (the sequential version does not fully utilize the processor).

The third reason is that concurrency provides a different method of conceptualizing program solutions to problems. Many problem domains lend themselves naturally to concurrency in much the same way that recursion is a natural way to design solutions to some problems. Also, many programs are written to simulate physical entities and activities. In many cases, the system being simulated includes more than one entity, and the entities do whatever they do simultaneously—for example, aircraft flying in a controlled airspace, relay stations in a communications network, and the various machines in a factory. Software that uses concurrency must be used to simulate such systems accurately.

The fourth reason for using concurrency is to program applications that are distributed over several machines, either locally or through the Internet. Many machines, for example cars, have more than one built-in computer, each of which is dedicated to some specific task. In many cases, these collections of computers must synchronize their program executions. Internet games are another example of software that is distributed over multiple processors.
Concurrency is now used in numerous everyday computing tasks. Web servers process document requests concurrently. Web browsers now use secondary core processors to run graphic processing and to interpret programming code embedded in documents. In every operating system there are many concurrent processes being executed at all times, managing resources, getting input from keyboards, displaying output from programs, and reading and writing external memory devices. In short, concurrency has become a ubiquitous part of computing.

13.2 Introduction to Subprogram-Level Concurrency

Before language support for concurrency can be considered, one must understand the underlying concepts of concurrency and the requirements for it to be useful. These topics are covered in this section.

13.2.1 Fundamental Concepts

A task is a unit of a program, similar to a subprogram, that can be in concurrent execution with other units of the same program. Each task in a program can support one thread of control. Tasks are sometimes called processes. In some languages, for example Java and C#, certain methods serve as tasks. Such methods are executed in objects called threads.

Three characteristics of tasks distinguish them from subprograms. First, a task may be implicitly started, whereas a subprogram must be explicitly called. Second, when a program unit invokes a task, in some cases it need not wait for the task to complete its execution before continuing its own. Third, when the execution of a task is completed, control may or may not return to the unit that started that execution.

Tasks fall into two general categories: heavyweight and lightweight. Simply stated, a heavyweight task executes in its own address space. Lightweight tasks all run in the same address space. It is easier to implement lightweight tasks than heavyweight tasks. Furthermore, lightweight tasks can be more efficient than heavyweight tasks, because less effort is required to manage their execution.

A task can communicate with other tasks through shared nonlocal variables, through message passing, or through parameters. If a task does not communicate with or affect the execution of any other task in the program in any way, it is said to be disjoint. Because tasks often work together to create simulations or solve problems and therefore are not disjoint, they must use some form of communication to either synchronize their executions or share data or both.

Synchronization is a mechanism that controls the order in which tasks execute. Two kinds of synchronization are required when tasks share data: cooperation and competition. Cooperation synchronization is required between task A and task B when task A must wait for task B to complete some specific activity before task A can begin or continue its execution. Competition synchronization is required between two tasks when both require the use of
some resource that cannot be simultaneously used. Specifically, if task A needs to access shared data location x while task B is accessing x, task A must wait for task B to complete its processing of x. So, for cooperation synchronization, tasks may need to wait for the completion of specific processing on which their correct operation depends, whereas for competition synchronization, tasks may need to wait for the completion of any other processing by any task currently occurring on specific shared data.

A simple form of cooperation synchronization can be illustrated by a common problem called the producer-consumer problem. This problem originated in the development of operating systems, in which one program unit produces some data value or resource and another uses it. Produced data are usually placed in a storage buffer by the producing unit and removed from that buffer by the consuming unit. The sequence of stores to and removals from the buffer must be synchronized. The consumer unit must not be allowed to take data from the buffer if the buffer is empty. Likewise, the producer unit cannot be allowed to place new data in the buffer if the buffer is full. This is a problem of cooperation synchronization because the users of the shared data structure must cooperate if the buffer is to be used correctly.

Competition synchronization prevents two tasks from accessing a shared data structure at exactly the same time—a situation that could destroy the integrity of that shared data. To provide competition synchronization, mutually exclusive access to the shared data must be guaranteed.

To clarify the competition problem, consider the following scenario: Suppose task A has the statement \( TOTAL += 1 \), where \( TOTAL \) is a shared integer variable. Furthermore, suppose task B has the statement \( TOTAL *= 2 \). Task A and task B could try to change \( TOTAL \) at the same time.

At the machine language level, each task may accomplish its operation on \( TOTAL \) with the following three-step process:

1. Fetch the value of \( TOTAL \).
2. Perform the arithmetic operation.
3. Put the new value back in \( TOTAL \).

Without competition synchronization, given the previously described operations performed by tasks A and B on \( TOTAL \), four different values could result, depending on the order of the steps of the operations. Assume \( TOTAL \) has the value 3 before either A or B attempts to modify it. If task A completes its operation before task B begins, the value will be 8, which is assumed here to be correct. But if both A and B fetch the value of \( TOTAL \) before either task puts its new value back, the result will be incorrect. If A puts its value back first, the value of \( TOTAL \) will be 6. This case is shown in Figure 13.1. If B puts its value back first, the value of \( TOTAL \) will be 4. Finally, if B completes its operation before task A begins, the value will be 7. A situation that leads to these problems is sometimes called a race condition, because two or more tasks are racing to use the shared resource and the behavior of the program depends on which task arrives first (and wins the race). The importance of competition synchronization should now be clear.
One general method for providing mutually exclusive access (to support competition synchronization) to a shared resource is to consider the resource to be something that a task can possess and allow only a single task to possess it at a time. To gain possession of a shared resource, a task must request it. Possession will be granted only when no other task has possession. While a task possesses a resource, all other tasks are prevented from having access to that resource. When a task is finished with a shared resource that it possesses, it must relinquish that resource so it can be made available to other tasks.

Three methods of providing for mutually exclusive access to a shared resource are semaphores, which are discussed in Section 13.3; monitors, which are discussed in Section 13.4; and message passing, which is discussed in Section 13.5.

Mechanisms for synchronization must be able to delay task execution. Synchronization imposes an order of execution on tasks that is enforced with these delays. To understand what happens to tasks through their lifetimes, we must consider how task execution is controlled. Regardless of whether a machine has a single processor or more than one, there is always the possibility of there being more tasks than there are processors. A run-time system program called a scheduler manages the sharing of processors among the tasks. If there were never any interruptions and tasks all had the same priority, the scheduler could simply give each task a time slice, such as 0.1 second, and when a task’s turn came, the scheduler could let it execute on a processor for that amount of time. Of course, there are several events that complicate this, for example, task delays for synchronization and for input or output operations. Because input and output operations are very slow relative to the processor’s speed, a task is not allowed to keep a processor while it waits for completion of such an operation.

Tasks can be in several different states:

1. New: A task is in the new state when it has been created but has not yet begun its execution.

2. Ready: A ready task is ready to run but is not currently running. Either it has not been given processor time by the scheduler, or it had run previously but was blocked in one of the ways described in Paragraph 4 of
this subsection. Tasks that are ready to run are stored in a queue that is often called the **task-ready queue**.

3. **Running**: A running task is one that is currently executing; that is, it has a processor and its code is being executed.

4. **Blocked**: A task that is blocked has been running, but that execution was interrupted by one of several different events, the most common of which is an input or output operation. In addition to input and output, some languages provide operations for the user program to specify that a task be blocked.

5. **Dead**: A dead task is no longer active in any sense. A task dies when its execution is completed or it is explicitly killed by the program.

A flow diagram of the states of a task is shown in Figure 13.2. One important issue in task execution is the following: How is a ready task chosen to move to the running state when the task currently running has become blocked or whose time slice has expired? Several different algorithms
have been used for making this choice, some based on specifiable priority levels. The algorithm that does the choosing is implemented in the scheduler.

Associated with the concurrent execution of tasks and the use of shared resources is the concept of liveness. In the environment of sequential programs, a program has the liveness characteristic if it continues to execute, eventually leading to completion. In more general terms, liveness means that if some event—say, program completion—is supposed to occur, it will occur, eventually. That is, progress is continually made. In a concurrent environment and with shared resources, the liveness of a task can cease to exist, meaning that the program cannot continue and thus will never terminate.

For example, suppose task $A$ and task $B$ both need the shared resources $X$ and $Y$ to complete their work. Furthermore, suppose that task $A$ gains possession of $X$ and task $B$ gains possession of $Y$. After some execution, task $A$ needs resource $Y$ to continue, so it requests $Y$ but must wait until $B$ releases it. Likewise, task $B$ requests $X$ but must wait until $A$ releases it. Neither relinquishes the resource it possesses, and as a result, both lose their liveness, guaranteeing that execution of the program will never complete normally. This particular kind of loss of liveness is called deadlock. Deadlock is a serious threat to the reliability of a program, and therefore its avoidance demands serious consideration in both language and program design.

We are now ready to discuss some of the linguistic mechanisms for providing concurrent unit control.

### 13.2.2 Language Design for Concurrency

In some cases, concurrency is implemented through libraries. Among these is OpenMP, an applications programming interface to support shared memory multiprocessor programming in C, C++, and Fortran on a variety of platforms. Our interest in this book, of course, is language support for concurrency. A number of languages have been designed to support concurrency, beginning with PL/I in the mid-1960s and including the contemporary languages Ada 95, Java, C#, F#, Python, and Ruby.¹

### 13.2.3 Design Issues

The most important design issues for language support for concurrency have already been discussed at length: competition and cooperation synchronization. In addition to these, there are several design issues of secondary importance. Prominent among them is how an application can influence task scheduling. Also, there are the issues of how and when tasks start and end their executions, and how and when they are created.

¹. In the cases of Python and Ruby, programs are interpreted, so there only can be logical concurrency. Even if the machine has multiple processors, these programs cannot make use of more than one.
Keep in mind that our discussion of concurrency is intentionally incomplete, and only the most important of the language design issues related to support for concurrency are discussed.

The following sections discuss three alternative approaches to the design issues for concurrency: semaphores, monitors, and message passing.

**13.3 Semaphores**

A semaphore is a simple mechanism that can be used to provide synchronization of tasks. Although semaphores are an early approach to providing synchronization, they are still used, both in contemporary languages and in library-based concurrency support systems. In the following paragraphs, we describe semaphores and discuss how they can be used for this purpose.

**13.3.1 Introduction**

In an effort to provide competition synchronization through mutually exclusive access to shared data structures, Edsger Dijkstra devised semaphores in 1965 (Dijkstra, 1968b). Semaphores can also be used to provide cooperation synchronization.

To provide limited access to a data structure, guards can be placed around the code that accesses the structure. A *guard* is a linguistic device that allows the guarded code to be executed only when a specified condition is true. So, a guard can be used to allow only one task to access a particular shared data structure at a time. A semaphore is an implementation of a guard. Specifically, a *semaphore* is a data structure that consists of an integer and a queue that stores task descriptors. A *task descriptor* is a data structure that stores all of the relevant information about the execution state of a task.

An integral part of a guard mechanism is a procedure for ensuring that all attempted executions of the guarded code eventually take place. The typical approach is to have requests for access that occur when access cannot be granted or stored in the task descriptor queue, from which they are later allowed to leave and execute the guarded code. This is the reason a semaphore must have both a counter and a task descriptor queue.

The only two operations provided for semaphores were originally named P and V by Dijkstra, after the two Dutch words *passeren* (to pass) and *vrygeren* (to release) (Andrews and Schneider, 1983). We will refer to these as *wait* and *release*, respectively, in the remainder of this section.

**13.3.2 Cooperation Synchronization**

Through much of this chapter, we use the example of a shared buffer used by producers and consumers to illustrate the different approaches to providing cooperation and competition synchronization. For cooperation synchronization, such a buffer must have some way of recording both the number
of empty positions and the number of filled positions in the buffer (to prevent buffer underflow and overflow). The counter component of a semaphore can be used for this purpose. One semaphore variable—for example, `emptyspots`—can use its counter to maintain the number of empty locations in a shared buffer used by producers and consumers, and another—say, `fullspots`—can use its counter to maintain the number of filled locations in the buffer. The queues of these semaphores can store the descriptors of tasks that have been forced to wait for access to the buffer. The queue of `emptyspots` can store producer tasks that are waiting for available positions in the buffer; the queue of `fullspots` can store consumer tasks waiting for values to be placed in the buffer.

Our example buffer is designed as an abstract data type in which all data enters the buffer through the subprogram `DEPOSIT`, and all data leaves the buffer through the subprogram `FETCH`. The `DEPOSIT` subprogram needs only to check with the `emptyspots` semaphore to see whether there are any empty positions. If there is at least one, it can proceed with the `DEPOSIT`, which must have the side effect of decrementing the counter of `emptyspots`. If the buffer is full, the caller to `DEPOSIT` must be made to wait in the `emptyspots` queue for an empty spot to become available. When the `DEPOSIT` is complete, the `DEPOSIT` subprogram increments the counter of the `fullspots` semaphore to indicate that there is one more filled location in the buffer.

The `FETCH` subprogram has the opposite sequence of `DEPOSIT`. It checks the `fullspots` semaphore to see whether the buffer contains at least one item. If it does, an item is removed and the `emptyspots` semaphore has its counter incremented by 1. If the buffer is empty, the calling task is put in the `fullspots` queue to wait until an item appears. When `FETCH` is finished, it must increment the counter of `emptyspots`.

The operations on semaphore types often are not direct—they are done through `wait` and `release` subprograms. Therefore, the `DEPOSIT` operation just described is actually accomplished in part by calls to `wait` and `release`. Note that `wait` and `release` must be able to access the task-ready queue.

The `wait` semaphore subprogram is used to test the counter of a given semaphore variable. If the value is greater than zero, the caller can carry out its operation. In this case, the counter value of the semaphore variable is decremented to indicate that there is now one fewer of whatever it counts. If the value of the counter is zero, the caller must be placed on the waiting queue of the semaphore variable, and the processor must be given to some other ready task.

The `release` semaphore subprogram is used by a task to allow some other task to have one of whatever the counter of the specified semaphore variable counts. If the queue of the specified semaphore variable is empty, which means no task is waiting, `release` increments its counter (to indicate there is one more of whatever is being controlled that is now available). If one or more tasks are waiting, `release` moves one of them from the semaphore queue to the ready queue.
The following are concise pseudocode descriptions of \texttt{wait} and \texttt{release}:

\begin{verbatim}
wait(aSemaphore)
  if aSemaphore’s counter > 0 then
    decrement aSemaphore’s counter
  else
    put the caller in aSemaphore’s queue
    attempt to transfer control to some ready task
    (if the task-ready queue is empty, deadlock occurs)
  end if

release(aSemaphore)
  if aSemaphore’s queue is empty (no task is waiting) then
    increment aSemaphore’s counter
  else
    put the calling task in the task-ready queue
    transfer control to a task from aSemaphore’s queue
  end
\end{verbatim}

We can now present an example program that implements cooperation synchronization for a shared buffer. In this case, the shared buffer stores integer values and is a logically circular structure. It is designed for use by possibly multiple producer and consumer tasks.

The following pseudocode shows the definition of the producer and consumer tasks. Two semaphores are used to ensure against buffer underflow or overflow, thus providing cooperation synchronization. Assume that the buffer has length \texttt{BUFLEN}, and the routines that actually manipulate it already exist as \texttt{FETCH} and \texttt{DEPOSIT}. Accesses to the counter of a semaphore are specified by dot notation. For example, if \texttt{fullspots} is a semaphore, its counter is referenced by \texttt{fullspots.count}.

\begin{verbatim}
semaphore fullspots, emptyspots;
fullspots.count = 0;
emptyspots.count = BUFLEN;
task producer;
  loop
    -- produce VALUE --
    wait(emptyspots);    \{ wait for a space \}
    DEPOSIT(VALUE);
    release(fullspots);  \{ increase filled spaces \}
  end loop;
end producer;

task consumer;
  loop
    wait(fullspots);    \{ make sure it is not empty \}
\end{verbatim}
The semaphore `fullspots` causes the `consumer` task to be queued to wait for a buffer entry if it is currently empty. The semaphore `emptyspots` causes the `producer` task to be queued to wait for an empty space in the buffer if it is currently full.

### 13.3.3 Competition Synchronization

Our buffer example does not provide competition synchronization. Access to the structure can be controlled with an additional semaphore. This semaphore need not count anything but can simply indicate with its counter whether the buffer is currently being used. The `wait` statement allows the access only if the semaphore’s counter has the value 1, which indicates that the shared buffer is not currently being accessed. If the semaphore’s counter has a value of 0, there is a current access taking place, and the task is placed in the queue of the semaphore. Notice that the semaphore’s counter must be initialized to 1. The queues of semaphores must always be initialized to empty before use of the queue can begin.

A semaphore that requires only a binary-valued counter, like the one used to provide competition synchronization in the following example, is called a **binary semaphore**.

The example pseudocode that follows illustrates the use of semaphores to provide both competition and cooperation synchronization for a concurrently accessed shared buffer. The `access` semaphore is used to ensure mutually exclusive access to the buffer. Remember that there may be more than one producer and more than one consumer.

```plaintext
semaphore access, fullspots, emptyspots;
access.count = 1;
fullspots.count = 0;
emptyspots.count = BUFLEN;

task producer;
  loop
    -- produce VALUE --
    wait(emptyspots); { wait for a space }
    wait(access);    { wait for access }
    DEPOSIT(VALUE);
    release(access);  { relinquish access }
    release(fullspots); { increase filled spaces }
  end loop;
end producer;
```
task consumer;
    loop
        wait(fullspots);       { make sure it is not empty }
        wait(access);         { wait for access }
        FETCH(VALUE);
        release(access);      { relinquish access }
        release(emptyspots);  { increase empty spaces }
        -- consume VALUE --
    end loop
end consumer;

A brief look at this example may lead one to believe that there is a problem. Specifically, suppose that while a task is waiting at the `wait(access)` call in `consumer`, another task takes the last value from the shared buffer. Fortunately, this cannot happen, because the `wait(fullspots)` reserves a value in the buffer for the task that calls it by decrementing the `fullspots` counter.

There is one crucial aspect of semaphores that thus far has not been discussed. Recall the earlier description of the problem of competition synchronization: Operations on shared data must not overlap. If a second operation begins while an earlier operation is still in progress, the shared data can become corrupted. A semaphore is itself a shared data object, so the operations on semaphores are also susceptible to the same problem. It is therefore essential that semaphore operations be uninterruptible. Many computers have uninterruptible instructions that were designed specifically for semaphore operations. If such instructions are not available, then using semaphores to provide competition synchronization is a serious problem with no simple solution.

### 13.3.4 Evaluation

Using semaphores to provide cooperation synchronization creates an unsafe programming environment. There is no way to check statically for the correctness of their use, which depends on the semantics of the program in which they appear. In the buffer example, leaving out the `wait(emptyspots)` statement of the `producer` task would result in buffer overflow. Leaving out the `wait(-fullspots)` statement of the `consumer` task would result in buffer underflow. Leaving out either of the releases would result in deadlock. These are cooperation synchronization failures.

The reliability problems that semaphores cause in providing cooperation synchronization also arise when using them for competition synchronization. Leaving out the `wait(access)` statement in either task can cause insecure access to the buffer. Leaving out the `release(access)` statement in either task results in deadlock. These are competition synchronization failures. Noting the danger in using semaphores, Per Brinch Hansen (1973) wrote, “The semaphore is an elegant synchronization tool for an ideal programmer who never makes mistakes.” Unfortunately, ideal programmers are rare.
13.4 Monitors

One solution to some of the problems of semaphores in a concurrent environment is to encapsulate shared data structures with their operations and hide their representations—that is, to make shared data structures abstract data types with some special restrictions. This solution can provide competition synchronization without semaphores by transferring responsibility for synchronization to the run-time system.

13.4.1 Introduction

When the concepts of data abstraction were being formulated, the people involved in that effort applied the same concepts to shared data in concurrent programming environments to produce monitors. According to Per Brinch Hansen (Brinch Hansen, 1977, p. xvi), Edsger Dijkstra suggested in 1971 that all synchronization operations on shared data be gathered into a single program unit. Brinch Hansen (1973) formalized this concept in the environment of operating systems. The following year, Hoare (1974) named these structures monitors.

The first programming language to incorporate monitors was Concurrent Pascal (Brinch Hansen, 1975). Modula (Wirth, 1977), CSP/k (Holt et al., 1978), and Mesa (Mitchell et al., 1979) also provide monitors. Among contemporary languages, monitors are supported by Ada, Java, and C#, all of which are discussed later in this chapter.

13.4.2 Competition Synchronization

One of the most important features of monitors is that shared data is resident in the monitor rather than in any of the client units. The programmer does not synchronize mutually exclusive access to shared data through the use of semaphores or other mechanisms. Because the access mechanisms are part of the monitor, implementation of a monitor can be made to guarantee synchronized access by allowing only one access at a time. Calls to monitor procedures are implicitly blocked and stored in a queue if the monitor is busy at the time of the call.

13.4.3 Cooperation Synchronization

Although mutually exclusive access to shared data is intrinsic with a monitor, cooperation between processes is still the task of the programmer. In particular, the programmer must guarantee that a shared buffer does not experience underflow or overflow. Different languages provide different ways of programming cooperation synchronization, all of which are related to semaphores.

A program containing four tasks and a monitor that provides synchronized access to a concurrently shared buffer is shown in Figure 13.3. In this figure,
the interface to the monitor is shown as the two boxes labeled `insert` and `remove` (for the insertion and removal of data). The monitor appears exactly like an abstract data type—a data structure with limited access— which is what a monitor is.

**13.4.4 Evaluation**

Monitors are a better way to provide competition synchronization than are semaphores, primarily because of the problems of semaphores, as discussed in Section 13.3. The cooperation synchronization is still a problem with monitors, as will be clear when Ada and Java implementations of monitors are discussed in the following sections.

Semaphores and monitors are equally powerful at expressing concurrency control—semaphores can be used to implement monitors and monitors can be used to implement semaphores.

Ada provides two ways to implement monitors. Ada 83 includes a general tasking model that can be used to support monitors. Ada 95 added a cleaner and more efficient way of constructing monitors, called `protected objects`. Both of these approaches use message passing as a basic model for supporting concurrency. The message-passing model allows concurrent units to be distributed, which monitors do not allow. Message passing is described in Section 13.5; Ada support for message passing is discussed in Section 13.6.

In Java, a monitor can be implemented in a class designed as an abstract data type, with the shared data being the type. Accesses to objects of the class are controlled by adding the `synchronized` modifier to the access methods. An example of a monitor for the shared buffer written in Java is given in Section 13.7.4.
C# has a predefined class, `Monitor`, which is designed for implementing monitors.

### 13.5 Message Passing

This section introduces the fundamental concept of message passing in concurrency. Note that this concept of message passing is unrelated to the message passing used in object-oriented programming to enact methods.

#### 13.5.1 Introduction

The first efforts to design languages that provide the capability for message passing among concurrent tasks were those of Brinch Hansen (1978) and Hoare (1978). These pioneer developers of message passing also developed a technique for handling the problem of what to do when multiple simultaneous requests were made by other tasks to communicate with a given task. It was decided that some form of nondeterminism was required to provide fairness in choosing which among those requests would be taken first. This fairness can be defined in various ways, but in general, it means that all requesters are provided an equal chance of communicating with a given task (assuming that every requester has the same priority). Nondeterministic constructs for statement-level control, called guarded commands, were introduced by Dijkstra (1975). Guarded commands are discussed in Chapter 8. Guarded commands are the basis of the construct designed for controlling message passing.

#### 13.5.2 The Concept of Synchronous Message Passing

Message passing can be either synchronous or asynchronous. Here, we describe synchronous message passing. The basic concept of synchronous message passing is that tasks are often busy, and when busy, they cannot be interrupted by other units. Suppose task A and task B are both in execution, and A wishes to send a message to B. Clearly, if B is busy, it is not desirable to allow another task to interrupt it. That would disrupt B’s current processing. Furthermore, messages usually cause associated processing in the receiver, which might not be sensible if other processing is incomplete. The alternative is to provide a linguistic mechanism that allows a task to specify to other tasks when it is ready to receive messages. This approach is somewhat like an executive who instructs his or her secretary to hold all incoming calls until another activity, perhaps an important conversation, is completed. Later, when the current conversation is complete, the executive tells the secretary that he or she is now willing to talk to one of the callers who has been placed on hold.

A task can be designed so that it can suspend its execution at some point, either because it is idle or because it needs information from another unit before it can continue. This is like a person who is waiting for an important call. In some cases, there is nothing else to do but sit and wait. However, if task A is
waiting for a message at the time task \( B \) sends that message, the message can be transmitted. This actual transmission of the message is called a **rendezvous**. Note that a rendezvous can occur only if both the sender and receiver want it to happen. During a rendezvous, the information of the message can be transmitted in either or both directions.

Both cooperation and competition synchronization of tasks can be conveniently handled with the message-passing model, as described in the following section.

## 13.6 Ada Support for Concurrency

This section describes the support for concurrency provided by Ada. Ada 83 supports only synchronous message passing.

### 13.6.1 Fundamentals

The Ada design for tasks is partially based on the work of Brinch Hansen and Hoare in that message passing is the design basis and nondeterminism is used to choose among the tasks that have sent messages.

The full Ada tasking model is complex, and the following discussion of it is limited. The focus here will be on the Ada version of the synchronous message-passing mechanism.

Ada tasks can be more active than monitors. Monitors are passive entities that provide management services for the shared data they store. They provide their services, though only when those services are requested. When used to manage shared data, Ada tasks can be thought of as managers that can reside with the resource they manage. They have several mechanisms, some deterministic and some nondeterministic, that allow them to choose among competing requests for access to their resources.

There are two parts to an Ada task—a specification part and a body part—both with the same name. The interface of a task is its entry points or locations where it can accept messages from other tasks. Because these entry points are part of its interface, it is natural that they be listed in the specification part of a task. Because a rendezvous can involve an exchange of information, messages can have parameters; therefore, task entry points must also allow parameters, which must also be described in the specification part. In appearance, a task specification is similar to the package specification for an abstract data type.

As an example of an Ada task specification, consider the following code, which includes a single entry point named `Entry_1`, which has an in-mode parameter:

```ada
task Task_Example is
  entry Entry_1(Item : in Integer);
end Task_Example;
```
A task body must include some syntactic form of the entry points that correspond to the entry clauses in that task’s specification part. In Ada, these task body entry points are specified by clauses that are introduced by the accept reserved word. An accept clause is defined as the range of statements beginning with the accept reserved word and ending with the matching end reserved word. accept clauses are themselves relatively simple, but other constructs in which they can be embedded can make their semantics complex. A simple accept clause has the following form:

```
accept entry_name (formal parameters) do
    ...
end entry_name;
```

The accept entry name matches the name in an entry clause in the associated task specification part. The optional parameters provide the means of communicating data between the caller and the called task. The statements between the do and the end define the operations that take place during the rendezvous. These statements are together called the accept clause body. During the actual rendezvous, the sender task is suspended.

Whenever an accept clause receives a message that it is not willing to accept, for whatever reason, the sender task must be suspended until the accept clause in the receiver task is ready to accept the message. Of course, the accept clause must also remember the sender tasks that have sent messages that were not accepted. For this purpose, each accept clause in a task has a queue associated with it that stores a list of other tasks that have unsuccessfully attempted to communicate with it.

The following is the skeletal body of the task whose specification was given previously:

```
task body Task_Example is
    begin
        loop
            accept Entry_1(Item : in Integer) do
                ...
            end Entry_1;
        end loop;
end Task_Example;
```

The accept clause of this task body is the implementation of the entry named Entry_1 in the task specification. If the execution of Task_Example begins and reaches the Entry_1 accept clause before any other task sends a message to Entry_1, Task_Example is suspended. If another task sends a message to Entry_1 while Task_Example is suspended at its accept, a rendezvous occurs and the accept clause body is executed. Then, because of the loop, execution proceeds back to the accept. If no other task has sent a message to Entry_1, execution is again suspended to wait for the next message.
A rendezvous can occur in two basic ways in this simple example. First, the receiver task, TaskExample, can be waiting for another task to send a message to the Entry_1 entry. When the message is sent, the rendezvous occurs. This is the situation described earlier. Second, the receiver task can be busy with one rendezvous, or with some other processing not associated with a rendezvous, when another task attempts to send a message to the same entry. In that case, the sender is suspended until the receiver is free to accept that message in a rendezvous. If several messages arrive while the receiver is busy, the senders are queued to wait their turn for a rendezvous.

The two rendezvous just described are illustrated with the timeline diagrams in Figure 13.4.

Tasks need not have entry points. Such tasks are called actor tasks because they do not wait for a rendezvous in order to do their work. Actor tasks can rendezvous with other tasks by sending them messages. In contrast to actor tasks, a task can have accept clauses but not have any code outside those accept clauses, so it can only react to other tasks. Such a task is called a server task.

An Ada task that sends a message to another task must know the entry name in that task. However, the opposite is not true: A task entry need not
know the name of the task from which it will accept messages. This asymmetry is in contrast to the design of the language known as CSP, or Communicating Sequential Processes (Hoare, 1978). In CSP, which also uses the message-passing model of concurrency, tasks accept messages only from explicitly named tasks. The disadvantage of this is that libraries of tasks cannot be built for general use.

The usual graphical method of describing a rendezvous in which task $A$ sends a message to task $B$ is shown in Figure 13.5.

---

Figure 13.5
Graphical representation of a rendezvous caused by a message sent from task $A$ to task $B$

Tasks are declared in the declaration part of a package, subprogram, or block. Statically created tasks\(^2\) begin executing at the same time as the statements in the code to which that declarative part is attached. For example, a task declared in a main program begins execution at the same time as the first statement in the code body of the main program. Task termination, which is a complex issue, is discussed later in this section.

Tasks may have any number of entries. The order in which the associated accept clauses appear in the task dictates the order in which messages can be accepted. If a task has more than one entry point and requires them to be able to receive messages in any order, the task uses a select statement to enclose the entries. For example, suppose a task models the activities of a bank teller, who must serve customers at a walk-up station inside the bank and also serve

\(^2\) Tasks can also be dynamically created, but such tasks are not covered here.
customers at a drive-up window. The following skeletal teller task illustrates a 
select construct:

\[
\text{task body Teller is}
\begin{align*}
\text{begin} \\
\text{loop} \\
\text{select} \\
\quad \text{accept Drive_Up(formal parameters) do} \\
\quad \quad \ldots \\
\quad \quad \text{end Drive_Up;} \\
\quad \ldots \\
\quad \text{or} \\
\quad \quad \text{accept Walk_Up(formal parameters) do} \\
\quad \quad \quad \ldots \\
\quad \quad \quad \text{end Walk_Up;} \\
\quad \quad \ldots \\
\quad \text{end select;} \\
\text{end loop;}
\end{align*}
\text{end Teller;}
\]

In this task, there are two accept clauses, \text{Walk_Up} and \text{Drive_Up}, each
of which has an associated queue. The action of the select, when it is
executed, is to examine the queues associated with the two accept clauses.
If one of the queues is empty, but the other contains at least one waiting
message (customer), the accept clause associated with the waiting message
or messages has a rendezvous with the task that sent the first message that
was received. If both accept clauses have empty queues, the select waits
until one of the entries is called. If both accept clauses have nonempty
queues, one of the accept clauses is nondeterministically chosen to have a
rendezvous with one of its callers. The loop forces the select statement to
be executed repeatedly, forever.

The end of the accept clause marks the end of the code that assigns or
references the formal parameters of the accept clause. The code, if there
is any, between an accept clause and the next or (or the end select, if the
accept clause is the last one in the select) is called the extended accept
clause. The extended accept clause is executed only after the associated
(immediately preceding) accept clause is executed. This execution of the
extended accept clause is not part of the rendezvous and can take place
concurrently with the execution of the calling task. The sender is suspended
during the rendezvous, but it is put back in the ready queue when the end of
the accept clause is reached. If an accept clause has no formal parameters,
the do-end is not required, and the accept clause can consist entirely of
an extended accept clause. Such an accept clause would be used exclu-
sively for synchronization. Extended accept clauses are illustrated in the
Buf_Task task in Section 13.6.3.
13.6.2 Cooperation Synchronization

Each accept clause can have a guard attached, in the form of a when clause, that can delay rendezvous. For example,

\[
\text{when not Full(Buffer) =>}
\begin{align*}
\text{accept Deposit(New_Value) do} \\
& \ldots \\
\text{end}
\end{align*}
\]

An accept clause with a when clause is either open or closed. If the Boolean expression of the when clause is currently true, that accept clause is called open; if the Boolean expression is false, the accept clause is called closed. An accept clause that does not have a guard is always open. An open accept clause is available for rendezvous; a closed accept clause cannot rendezvous.

Suppose there are several guarded accept clauses in a select clause. Such a select clause is usually placed in an infinite loop. The loop causes the select clause to be executed repeatedly, with each when clause evaluated on each repetition. Each repetition causes a list of open accept clauses to be constructed. If exactly one of the open clauses has a nonempty queue, a message from that queue is taken and a rendezvous takes place. If more than one of the open accept clauses has nonempty queues, one queue is chosen nondeterministically, a message is taken from that queue, and a rendezvous takes place. If the queues of all open clauses are empty, the task waits for a message to arrive at one of those accept clauses, at which time a rendezvous will occur. If a select is executed and every accept clause is closed, a run-time exception or error results. This possibility can be avoided either by making sure one of the when clauses is always true or by adding an else clause in the select. An else clause can include any sequence of statements, except an accept clause.

A select clause may have a special statement, terminate, that is selected only when it is open and no other accept clause is open. A terminate clause, when selected, means that the task is finished with its job but is not yet terminated. Task termination is discussed later in this section.

13.6.3 Competition Synchronization

The features described so far provide for cooperation synchronization and communication among tasks. Next, we discuss how mutually exclusive access to shared data structures can be enforced in Ada.

If access to a data structure is to be controlled by a task, then mutually exclusive access can be achieved by declaring the data structure within a task. The semantics of task execution usually guarantees mutually exclusive access to the structure, because only one accept clause in the task can be active at a given time. The only exceptions to this occur when tasks are nested in procedures or other tasks. For example, if a task that defines a shared data structure
has a nested task, that nested task can also access the shared structure, which could destroy the integrity of the data. Thus, tasks that are meant to control access to a shared data structure should not define tasks.

The following is an example of an Ada task that implements a monitor for a buffer. The buffer behaves very much like the buffer in Section 13.3, in which synchronization is controlled with semaphores.

```ada
task Buf_Task is
  entry Deposit(Item : in Integer);
  entry Fetch(Item : out Integer);
end Buf_Task;

task body Buf_Task is
  Bufsize : constant Integer := 100;
  Buf   : array (1..Bufsize) of Integer;
  Filled : Integer range 0..Bufsize := 0;
  Next_In,
  Next_Out : Integer range 1..Bufsize := 1;
begin
  loop
    select
      when Filled < Bufsize =>
        accept Deposit(Item : in Integer) do
          Buf(Next_In) := Item;
        end Deposit;
        Next_In := (Next_In mod Bufsize) + 1;
        Filled := Filled + 1;
      or
      when Filled > 0 =>
        accept Fetch(Item : out Integer) do
          Item := Buf(Next_Out);
        end Fetch;
        Next_Out := (Next_Out mod Bufsize) + 1;
        Filled := Filled - 1;
    end select;
  end loop;
end Buf_Task;
```

In this example, both `accept` clauses are extended. These extended clauses can be executed concurrently with the tasks that called the associated `accept` clauses.

The tasks for a producer and a consumer that could use `Buf_Task` have the following form:

```ada
task Producer;
task Consumer;
task body Producer is
  New_Value : Integer;
beg
loop
  -- produce New_Value --
  Buf_Task.Deposit(New_Value);
end loop;
end Producer;

task body Consumer is
  Stored_Value : Integer;
begin
  loop
    Buf_Task.Fetch(Stored_Value);
    -- consume Stored_Value --
  end loop;
end Consumer;

13.6.4 Protected Objects

As we have seen, access to shared data can be controlled by enclosing the data in a task and allowing access only through task entries, which implicitly provide competition synchronization. One problem with this method is that it is difficult to implement the rendezvous mechanism efficiently. Ada 95 protected objects provide an alternative method of providing competition synchronization that need not involve the rendezvous mechanism.

A protected object is not a task; it is more like a monitor, as described in Section 13.4. Protected objects can be accessed either by protected subprograms or by entries that are syntactically similar to the `accept` clauses in tasks. The protected subprograms can be either protected procedures, which provide mutually exclusive read-write access to the data of the protected object, or protected functions, which provide concurrent read-only access to that data. Entries differ from protected subprograms in that they can have guards.

Within the body of a protected procedure, the current instance of the enclosing protected unit is defined to be a variable; within the body of a protected function, the current instance of the enclosing protected unit is defined to be a constant, which allows concurrent read-only access.

Entry calls to a protected object provide synchronous communication with one or more tasks using the same protected object. These entry calls provide access similar to that provided to the data enclosed in a task.

The buffer problem that is solved with a task in the previous subsection can be more simply solved with a protected object. Note that this example does not include protected subprograms.

protected Buffer is
  entry Deposit(Item : in Integer);
  entry Fetch(Item : out Integer);

3. Entries in protected object bodies use the reserved word `entry`, rather than the `accept` used in task bodies.
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private
BuFSIZE : constant Integer := 100;
Buf   : array (1..BuFSIZE) of Integer;
Filled : Integer range 0..BuFSIZE := 0;
Next_In,
Next_Out : Integer range 1..BuFSIZE := 1;
end Buffer;

protected body Buffer is
entry Deposit(Item : in Integer) when Filled < BuFSIZE is
begin
Buf(Next_In) := Item;
Next_In := (Next_In mod BuFSIZE) + 1;
Filled := Filled + 1;
end Deposit;
entry Fetch(Item : out Integer) when Filled > 0 is
begin Item := Buf(Next_Out);
Next_Out := (Next_Out mod BuFSIZE) + 1;
Filled := Filled - 1;
end Fetch;
end Buffer;

13.6.5  Evaluation

Using the general message-passing model of concurrency to construct monitors is like using Ada packages to support abstract data types—both are tools that are more general than is necessary. Protected objects are a better way to provide synchronized access to shared data.

In the absence of distributed processors with independent memories, the choice between monitors and tasks with message passing as a means of implementing synchronized access to shared data in a concurrent environment is somewhat a matter of taste. However, in the case of Ada, protected objects are clearly better than tasks for supporting concurrent access to shared data. Not only is the code simpler; it is also much more efficient.

For distributed systems, message passing is a better model for concurrency, because it naturally supports the concept of separate processes executing in parallel on separate processors.

13.7  Java Threads

The concurrent units in Java are methods named run, whose code can be in concurrent execution with other such methods (of other objects) and with the main method. The process in which the run methods execute is called a thread. Java’s threads are lightweight tasks, which means that they all run in the same address space. This is different from Ada tasks, which are heavyweight threads
Java Threads

(they run in their own address spaces).\(^4\) One important result of this difference is that threads require far less overhead than Ada’s tasks.

There are two ways to define a class with a run method. One of these is to define a subclass of the predefined class Thread and override its run method. However, if the new subclass has a necessary natural parent, then defining it as a subclass of Thread obviously will not work. In these situations, we define a subclass that inherits from its natural parent and implements the Runnable interface. Runnable provides the run method protocol, so any class that implements Runnable must define run. An object of the class that implements Runnable is passed to the Thread constructor. So, this approach still requires a Thread object, as will be seen in the example in Section 13.7.5.

In Ada, tasks can be either actors or servers and tasks communicate with each other through accept clauses. Java run methods are all actors and there is no mechanism for them to communicate with each other, except for the join method (see Section 13.7.1) and through shared data.

Java threads is a complex topic—this section only provides an introduction to its simplest but most useful parts.

### 13.7.1 The Thread Class

The Thread class is not the natural parent of any other classes. It provides some services for its subclasses, but it is not related in any natural way to their computational purposes. Thread is the only class available for creating concurrent Java programs. As previously stated, Section 13.7.5 will briefly discuss the use of the Runnable interface.

The Thread class includes five constructors and a collection of methods and constants. The run method, which describes the actions of the thread, is always overridden by subclasses of Thread. The start method of Thread starts its thread as a concurrent unit by calling its run method.\(^5\) The call to start is unusual in that control returns immediately to the caller, which then continues its execution, in parallel with the newly started run method.

Following is a skeletal subclass of Thread and a code fragment that creates an object of the subclass and starts the run method’s execution in the new thread:

```java
class MyThread extends Thread {
    public void run() { . . . }
}
.
.
Thread myTh = new MyThread();
myTh.start();
```

---

4. Actually, although Ada tasks behave as if they were heavyweight tasks, in some cases, they are now implemented as threads. This is sometimes done using libraries, such as the IBM Rational Apex Native POSIX Threading Library.

5. Calling the run method directly does not always work, because initialization that is sometimes required is included in the start method.
When a Java application program begins execution, a new thread is created (in which the main method will run) and main is called. Therefore, all Java application programs run in threads.

When a program has multiple threads, a scheduler must determine which thread or threads will run at any given time. In many cases, there is only a single processor available, so only one thread actually runs at a time. It is difficult to give a precise description of how the Java scheduler works, because the different implementations (Solaris, Windows, and so on) do not necessarily schedule threads in exactly the same way. Typically, however, the scheduler gives equal-size time slices to each ready thread in round-robin fashion, assuming all of these threads have the same priority. Section 13.7.2 describes how different priorities can be given to different threads.

The Thread class provides several methods for controlling the execution of threads. The yield method, which takes no parameters, is a request from the running thread to surrender the processor voluntarily. The thread is put immediately in the task-ready queue, making it ready to run. The scheduler then chooses the highest-priority thread from the task-ready queue. If there are no other ready threads with priority higher than the one that just yielded the processor, it may also be the next thread to get the processor.

The sleep method has a single parameter, which is the integer number of milliseconds that the caller of sleep wants the thread to be blocked. After the specified number of milliseconds has passed, the thread will be put in the task-ready queue. Because there is no way to know how long a thread will be in the task-ready queue before it runs, the parameter to sleep is the minimum amount of time the thread will not be in execution. The sleep method can throw an InterruptedException, which must be handled in the method that calls sleep. Exceptions are described in detail in Chapter 14.

The join method is used to force a method to delay its execution until the run method of another thread has completed its execution. join is used when the processing of a method cannot continue until the work of the other thread is complete. For example, we might have the following run method:

```java
public void run() {
    Thread myTh = new Thread();
    myTh.start();
    // do part of the computation of this thread
    myTh.join(); // Wait for myTh to complete
    // do the rest of the computation of this thread
}
```

The join method puts the thread that calls it in the blocked state, which can be ended only by the completion of the thread on which join was called. If that thread happens to be blocked, there is the possibility of deadlock. To prevent

---

6. The yield method is actually defined to be a “suggestion” to the scheduler, which it may or may not follow (though it usually does).
this, `join` can be called with a parameter, which is the time limit in milliseconds of how long the calling thread will wait for the called thread to complete. For example, the following call to `join` will cause the calling thread to wait two seconds for `myTh` to complete. If it has not completed its execution after two seconds have passed, the calling thread is put back in the ready queue, which means that it will continue its execution as soon as it is scheduled.

```java
myTh.join(2000);
```

Early versions of Java included three more `Thread` methods: `stop`, `suspend`, and `resume`. All three of these have been deprecated because of safety problems. The `stop` method is sometimes overridden with a simple method that destroys the thread by setting its reference variable to `null`.

The normal way a `run` method ends its execution is by reaching the end of its code. However, in many cases, threads run until told to terminate. Regarding this, there is the question of how a thread can determine whether it should continue or end. The `interrupt` method is one way to communicate to a thread that it should stop. This method does not stop the thread; rather, it sends the thread a message that actually just sets a bit in the thread object, which can be checked by the thread. The bit is checked with the predicate method, `isInterrupted`. This is not a complete solution, because the thread one is attempting to interrupt may be sleeping or waiting at the time the `interrupt` method is called, which means that it will not be checking to see if it has been interrupted. For these situations, the `interrupt` method also throws an exception, `InterruptedException`, which also causes the thread to awaken (from sleeping or waiting). So, a thread can periodically check to see whether it has been interrupted and if so, whether it can terminate. The thread cannot miss the interrupt, because if it was asleep or waiting when the interrupt occurred, it will be awakened by the interrupt. Actually, there are more details to the actions and uses of `interrupt`, but they are not covered here (Arnold et al., 2006).

### 13.7.2 Priorities

The priorities of threads need not all be the same. A thread's default priority initially is the same as the thread that created it. If `main` creates a thread, its default priority is the constant `NORM_PRIORITY`, which is usually 5. `Thread` defines two other priority constants, `MAX_PRIORITY` and `MIN_PRIORITY`, whose values are usually 10 and 1, respectively. The priority of a thread can be changed with the method `setPriority`. The new priority can be any of the predefined constants or any other number between `MIN_PRIORITY` and `MAX_PRIORITY`. The `getPriority` method returns the current priority of a thread. The priority constants are defined in `Thread`.

---

7. The number of priorities is implementation dependent, so there may be fewer or more than 10 levels in some implementations.
When there are threads with different priorities, the scheduler’s behavior is controlled by those priorities. When the executing thread is blocked or killed or the time slice for it expires, the scheduler chooses the thread from the task-ready queue that has the highest priority. A thread with lower priority will run only if one of higher priority is not in the task-ready queue when the opportunity arises.

### 13.7.3 Semaphores

The `java.util.concurrent.Semaphore` package defines the `Semaphore` class. Objects of this class implement counting semaphores. A counting semaphore has a counter, but no queue for storing thread descriptors. The `Semaphore` class defines two methods, `acquire` and `release`, which correspond to the `wait` and `release` operations described in Section 13.3.

The basic constructor for `Semaphore` takes one integer parameter, which initializes the semaphore’s counter. For example, the following could be used to initialize the `fullspots` and `emptyspots` semaphores for the buffer example of Section 13.3.2:

```java
fullspots = new Semaphore(0);
emptyspots = new Semaphore(BUFLEN);
```

The deposit operation of the producer method would appear as follows:

```java
emptyspots.acquire();
deposit(value);
fullspots.release();
```

Likewise, the fetch operation of the consumer method would appear as follows:

```java
fullspots.acquire();
fetch(value);
emptyspots.release();
```

The `deposit` and `fetch` methods could use the approach used in Section 13.7.4 to provide the competition synchronization required for the accesses to the buffer.

### 13.7.4 Competition Synchronization

Java methods (but not constructors) can be specified to be `synchronized`. A synchronized method called through a specific object must complete its execution before any other synchronized method can run on that object. Competition synchronization on an object is implemented by specifying that the methods that access shared data are synchronized. The synchronized mechanism is implemented as follows: Every Java object has a lock. Synchronized methods must acquire the lock of the object before they are allowed to execute, which prevents other synchronized methods from executing on the object.
during that time. A synchronized method releases the lock on the object on which it runs when it completes its execution, even if that completion is due to an exception. Consider the following skeletal class definition:

class ManageBuf {
    private int [100] buf;
    . . .
    public synchronized void deposit(int item) { . . . }
    public synchronized int fetch() { . . . }
    . . .
}

The two methods defined in ManageBuf are both defined to be synchronized, which prevents them from interfering with each other while executing on the same object, when they are called by separate threads.

An object whose methods are all synchronized is effectively a monitor. Note that an object may have one or more synchronized methods, as well as one or more unsynchronized methods. An unsynchronized method can run on an object at anytime, even during the execution of a synchronized method.

In some cases, the number of statements that deal with the shared data structure is significantly less than the number of other statements in the method in which it resides. In these cases, it is better to synchronize the code segment that changes the shared data structure rather than the whole method. This can be done with a so-called synchronized statement, whose general form is as follows:

```java
synchronized (expression){
    statements
}
```

The expression in this code must evaluate to an object and the statement can be a single statement or a compound statement. The object is locked during execution of the statement or compound statement, so the statement or compound statement is executed exactly as if it were the body of a synchronized method.

An object that has synchronized methods defined for it must have a queue associated with it that stores the synchronized methods that have attempted to execute on it while it was being operated upon by another synchronized method. Actually, every object has a queue called the intrinsic condition queue. These queues are implicitly supplied. When a synchronized method completes its execution on an object, a method that is waiting in the object’s intrinsic condition queue, if there is such a method, is put in the task-ready queue.

### 13.7.5 Cooperation Synchronization

Cooperation synchronization in Java is implemented with the `wait`, `notify`, and `notifyAll` methods, all of which are defined in `Object`, the root class of all Java classes. All classes except `Object` inherit these methods. Every
object has a wait list of all of the threads that have called \texttt{wait} on the object. The \texttt{notify} method is called to tell one waiting thread that an event that it may have been waiting for has occurred. The specific thread that is awakened by \texttt{notify} cannot be determined, because the Java Virtual Machine (JVM) chooses one from the wait list of the thread object at random. Because of this, along with the fact that the waiting threads may be waiting for different conditions, the \texttt{notifyAll} method is often used, rather than \texttt{notify}. The \texttt{notifyAll} method awakens all of the threads on the object’s wait list by putting them in the task-ready queue.

The methods \texttt{wait}, \texttt{notify}, and \texttt{notifyAll} can be called only from within a synchronized method, because they use the lock placed on an object by such a method. The call to \texttt{wait} is always put in a \texttt{while} loop that is controlled by the condition for which the method is waiting. The \texttt{while} loop is necessary because the \texttt{notify} or \texttt{notifyAll} that awakened the thread may have been called because of a change in a condition other than the one for which the thread was waiting. If it was a call to \texttt{notifyAll}, there is even a smaller chance that the condition being waited for is now true. Because of the use of \texttt{notifyAll}, some other thread may have changed the condition to false since it was last tested.

The \texttt{wait} method can throw \texttt{InterruptedException}, which is a descendant of \texttt{Exception}. Java’s exception handling is discussed in Chapter 14. Therefore, any code that calls \texttt{wait} must also catch \texttt{InterruptedException}.

Assuming the condition being waited for is called \texttt{theCondition}, the conventional way to use \texttt{wait} is as follows:

```java
try {
    while (!theCondition)
        wait();
    -- Do whatever is needed after theCondition comes true
} catch (InterruptedException myProblem) { . . . }
```

The following program implements a circular queue for storing \texttt{int} values. It illustrates both cooperation and competition synchronization.

```java
// Queue
// This class implements a circular queue for storing int
// values. It includes a constructor for allocating and
// initializing the queue to a specified size. It has
// synchronized methods for inserting values into and
// removing values from the queue.

class Queue {
    private int [] que;
    private int nextIn,
        nextOut,
        filled,
        queSize;
```
public Queue(int size) {
    que = new int [size];
    filled = 0;
    nextIn = 1;
    nextOut = 1;
    queSize = size;
}  /** end of Queue constructor

public synchronized void deposit (int item)
    throws InterruptedException {
    try {
        while (filled == queSize)
            wait();
        que [nextIn] = item;
        nextIn = (nextIn % queSize) + 1;
        filled++;
        notifyAll();
    }  /** end of try clause
    catch(InterruptedException e) {}
}  /** end of deposit method

public synchronized int fetch()
    throws InterruptedException {
    int item = 0;
    try {
        while (filled == 0)
            wait();
        item = que [nextOut];
        nextOut = (nextOut % queSize) + 1;
        filled--;
        notifyAll();
    }  /** end of try clause
    catch(InterruptedException e) {}
    return item;
}  /** end of fetch method
}  /** end of Queue class

Notice that the exception handler (catch) does nothing here.

Classes to define producer and consumer objects that could use the Queue class can be defined as follows:

class Producer extends Thread {
    private Queue buffer;
    public Producer(Queue que) {
        buffer = que;
    }
    public void run() {
        int new_item;
        while (true) {
            /** Create a new_item
class Consumer extends Thread {
    private Queue buffer;
    public Consumer(Queue que) {
        buffer = que;
    }
    public void run() {
        int stored_item;
        while (true) {
            stored_item = buffer.fetch();
            //-- Consume the stored_item
        }
    }
}

The following code creates a Queue object, and a Producer and a Consumer object, both attached to the Queue object, and starts their execution:

Queue buff1 = new Queue(100);
Producer producer1 = new Producer(buff1);
Consumer consumer1 = new Consumer(buff1);
producer1.start();
consumer1.start();

We could define one or both of the Producer and the Consumer as implementations of the Runnable interface rather than as subclasses of Thread. The only difference is in the first line, which would now appear as follows:

class Producer implements Runnable {
    . . .
}

To create and run an object of such a class, it is still necessary to create a Thread object that is connected to the object. This is illustrated in the following code:

Producer producer1 = new Producer(buff1);
Thread producerThread = new Thread(producer1);
producerThread.start();

Note that the buffer object is passed to the Producer constructor and the Producers object is passed to the Thread constructor.
13.7.6 Nonblocking Synchronization

Java includes some classes for controlling accesses to certain variables that do not include blocking or waiting. The java.util.concurrent.atomic package defines classes that allow certain nonblocking synchronized access to int, long, and boolean primitive type variables, as well as references and arrays. For example, the AtomicInteger class defines getter and setter methods, as well as methods for add, increment, and decrement operations. These operations are all atomic; that is, they cannot be interrupted, so locks are not required to guarantee the integrity of the values of the affected variables in a multithreaded program. This is fine-grained synchronization—just a single variable. Most machines now have atomic instructions for these operations on int and long types, so they are often easy to implement (implicit locks are not required).

The advantage of nonblocking synchronization is efficiency. A nonblocking access that does not occur during contention will be no slower, and usually faster than one that uses synchronized. A nonblocking access that occurs during contention definitely will be faster than one that uses synchronized, because the latter will require suspension and rescheduling of threads.

13.7.7 Explicit Locks

Java 5.0 introduced explicit locks as an alternative to synchronized method and blocks, which provide implicit locks. The Lock interface declares the lock, unlock, and tryLock methods. The predefined ReentrantLock class implements the Lock interface. To lock a block of code, the following idiom can be used:

```java
Lock lock = new ReentrantLock();
...
Lock.lock();
try {
    // The code that accesses the shared data
} finally {
    Lock.unlock();
}
```

This skeletal code creates a Lock object and calls the lock method on the Lock object. Then, it uses a try block to enclose the critical code. The call to unlock is in a finally clause to guarantee the lock is released, regardless of what happens in the try block.

There are at least two situations in which explicit locks are used rather than implicit locks: First, if the application needs to try to acquire a lock but cannot wait forever for it, the Lock interface includes a method, tryLock, that takes a time limit parameter. If the lock is not acquired within the time limit, execution continues at the statement following the call to tryLock. Second,
explicit locks are used when it is not convenient to have the lock-unlock pairs block structured. Implicit locks are always unlocked at the end of the compound statement in which they are locked. Explicit locks can be unlocked anywhere in the code, regardless of the structure of the program.

One danger of using explicit locks (and is not the case with using implicit locks) is that of omitting the unlock. Implicit locks are implicitly unlocked at the end of the locked block. However, explicit locks stay locked until explicitly unlocked, which can potentially be never.

As stated previously, each object has an intrinsic condition queue, which stores threads waiting for a condition on the object. The wait, notify, and notifyAll methods are the API for an intrinsic condition queue. Because each object can have just one condition queue, a queue may have threads in it waiting for different conditions. For example, the queue for our buffer example Queue can have threads waiting for either of two conditions (filled == queSize or filled == 0). That is the reason why the buffer uses notifyAll. (If it used notify, only one thread would be awakened, and it might be one that was waiting for a different condition than the one that actually became true.) However, notifyAll is expensive to use, because it awakens all threads waiting on an object and all must check their condition to determine which one runs. Furthermore, to check their condition, they must first acquire the lock on the object.

An alternative to using the intrinsic condition queue is the Condition interface, which uses a condition queue associated with a Lock object. It also declares alternatives to wait, notify, and notifyAll named await, signal, and signalAll. There can be any number of Condition objects with one Lock object. With Condition, signal, rather than signalAll, can be used, which is both easier to understand and more efficient, in part because it results in fewer context switches.

### 13.7.8 Evaluation

Java’s support for concurrency is relatively simple but effective. All Java run methods are actor tasks and there is no mechanism for communication, except through shared data, as there is among Ada tasks. Because they are heavyweight threads, Ada’s tasks easily can be distributed to different processors; in particular, different processors with different memories, which could be on different computers in different places. These kinds of systems are not possible with Java’s threads.

### 13.8 C# Threads

Although C#’s threads are loosely based on those of Java, there are significant differences. Following is a brief overview of C#’s threads.
13.8.1 Basic Thread Operations

Rather than just methods named `run`, as in Java, any C# method can run in its own thread. When C# threads are created, they are associated with an instance of a predefined delegate, `ThreadStart`. When execution of a thread is started, its delegate has the address of the method it is supposed to run. So, execution of a thread is controlled through its associated delegate.

A C# thread is created by creating a `Thread` object. The `Thread` constructor must be sent an instantiation of `ThreadStart`, to which must be sent the name of the method that is to run in the thread. For example, we might have

```csharp
public void MyRun1() { . . . }

Thread myThread = new Thread(new ThreadStart(MyRun1));
```

In this example, we create a thread named `myThread`, whose delegate points to the method `MyRun1`. So, when the thread begins execution it calls the method whose address is in its delegate. In this example, `myThread` is the delegate and `MyRun1` is the method.

As with Java, in C# there are two categories of threads: actors and servers. Actor threads are not called specifically; rather, they are started. Also, the methods that they execute do not take parameters or return values. As with Java, creating a thread does not start its concurrent execution. For actor threads, execution must be requested through a method of the `Thread` class, in this case named `Start`, as in

```csharp
myThread.Start();
```

As in Java, a thread can be made to wait for another thread to finish its execution before continuing, using the similarly named method `Join`. For example, suppose thread `A` has the following call:

```csharp
B.Join();
```

Thread `A` will be blocked until thread `B` exits.

The `Join` method can take an `int` parameter, which specifies a time limit in milliseconds that the caller will wait for the thread to finish.

A thread can be suspended for a specified amount of time with `Sleep`, which is a public static method of `Thread`. The parameter to `Sleep` is an integer number of milliseconds. Unlike its Java relative, C#’s `Sleep` does not raise any exceptions, so it need not be called in a `try` block.

A thread can be terminated with the `Abort` method, although it does not literally kill the thread. Instead, it throws `ThreadAbortException`, which the thread can catch. When the thread catches this exception, it usually deallocates any resources it allocated, and then ends (by getting to the end of its code).

A server thread runs only when called through its delegate. These threads are called servers because they provide some service when it is requested. Server
threads are more interesting than actor threads because they usually interact with other threads and often must have their execution synchronized with other threads.

Recall from Chapter 9, that any C# method can be called indirectly through a delegate. Such calls can be made by treating the delegate object as if it were the name of the method. This was actually an abbreviation for a call to a delegate method named Invoke. So, if a delegate object’s name is chgfun1 and the method it references takes one int parameter, we could call that method with either of the following statements:

```
chgfun1(7);
chgfun1.Invoke(7);
```

These calls are synchronous; that is, when the method is called, the caller is blocked until the method completes its execution. C# also supports asynchronous calls to methods that execute in threads. When a thread is called asynchronously, the called thread and the caller thread execute concurrently, because the caller is not blocked during the execution of the called thread.

A thread is called asynchronously through the delegate instance method BeginInvoke, to which are sent the parameters for the method of the delegate, along with two additional parameters, one of type AsyncCallback and the other of type object. BeginInvoke returns an object that implements the IAsyncResult interface. The delegate class also defines the EndInvoke instance method, which takes one parameter of type IAsyncResult and returns the same type that is returned by the method encapsulated in the delegate object. To call a thread asynchronously, we call it with BeginInvoke. For now, we will use null for the last two parameters. Suppose we have the following method declaration and thread definition:

```
public float MyMethod1(int x);
...  
Thread myThread = new Thread(new ThreadStart(MyMethod1));
```

The following statement calls MyMethod asynchronously:

```
IAsyncResult result = myThread.BeginInvoke(10, null, null);
```

The return value of the called thread is fetched with EndInvoke method, which takes as its parameter the object (of type IAsyncResult) returned by BeginInvoke. EndInvoke returns the return value of the called thread. For example, to get the float result of the call to MyMethod, we would use the following statement:

```
float returnValue = EndInvoke(result);
```

If the caller must continue some work while the called thread executes, it must have a way to determine when the called thread is finished. For this,
the IAsyncResult interface defines the IsCompleted property. While the called thread is executing, the caller can include code it can execute in a while loop that depends on IsCompleted. For example, we could have the following:

```csharp
IAsyncResult result = myThread.BeginInvoke(10, null, null);
while(!result.IsCompleted) {
    // Do some computation
}
```

This is an effective way to accomplish something in the calling thread while waiting for the called thread to complete its work. However, if the amount of computation in the while loop is relatively small, this is an inefficient way to use that time (because of the time required to test IsCompleted). An alternative is to give the called thread a delegate with the address of a callback method and have it call that method when it is finished. The delegate is sent as the second last parameter to BeginInvoke. For example, consider the following call to BeginInvoke:

```csharp
IAsyncResult result = myThread.BeginInvoke(10,
    new AsyncCallback(MyMethodComplete), null);
```

The callback method is defined in the caller. Such methods often simply set a Boolean variable, for example named isDone, to true. No matter how long the called thread takes, the callback method is called only once.

### 13.8.2 Synchronizing Threads

There are three different ways that C# threads can be synchronized: the Interlocked class, the Monitor class from the System.Threading namespace, and the lock statement. Each of these mechanisms is designed for a specific need. The Interlocked class is used when the only operations that need to be synchronized are the incrementing and decrementing of an integer. These operations are done atomically with the two methods of Interlocked, Increment and Decrement, both of which take a reference to an integer as the parameter. For example, to increment a shared integer named counter in a thread, we could use

```csharp
Interlocked.Increment(ref counter);
```

The lock statement is used to mark a critical section of code in a thread. The syntax of this is as follows:

```csharp
lock(token) {
    // The critical section
}
```
If the code to be synchronized is in a private instance method, the token is the current object, so `this` is used as the token for `lock`. If the code to be synchronized is in a public instance method, a new instance of `object` is created (in the class of the method with the code to be synchronized) and a reference to it is used as the token for `lock`.

The `Monitor` class defines five methods, `Enter`, `Wait`, `Pulse`, `PulseAll`, and `Exit`, which can be used to provide more control of the synchronization of threads. The `Enter` method, which takes an object reference as its parameter, marks the beginning of synchronization of the thread on that object. The `Wait` method suspends execution of the thread and instructs the Common Language Runtime (CLR) of .NET that this thread wants to resume its execution the next time there is an opportunity. The `Pulse` method, which also takes an object reference as its parameter, notifies one waiting thread that it now has a chance to run again. `PulseAll` is similar to Java's `notifyAll`. Threads that have been waiting are run in the order in which they called the `Wait` method. The `Exit` method ends the critical section of the thread.

The `lock` statement is compiled into a monitor, so `lock` is shorthand for a monitor. A monitor is used when the additional control (for example, with `Wait` and `PulseAll`) is needed.

.NET 4.0 added a collection of generic concurrent data structures, including structures for queues, stacks, and bags. These new classes are thread safe, meaning that they can be used in a multithreaded program without requiring the programmer to worry about competition synchronization. The `System.Collections.Concurrent` namespace defines these classes, whose names are `ConcurrentQueue<T>`, `ConcurrentStack<T>`, and `ConcurrentBag<T>`. So, our producer-consumer queue program could be written in C# using a `ConcurrentQueue<T>` for the data structure and there would be no need to program the competition synchronization for it. Because these concurrent collections are defined in .NET, they are also available in all of the other .NET languages.

### 13.8.3 Evaluation

C#'s threads are a slight improvement over those of its predecessor, Java. For one thing, any method can be run in its own thread. Recall that in Java, only methods named `run` can run in their own threads. Java supports actor threads only, but C# supports both actor and server threads. Thread termination is also cleaner with C# (calling a method (`Abort`) is more elegant than setting the thread's pointer to `null`). Synchronization of thread execution is more sophisticated in C#, because C# has several different mechanisms, each for a specific application. Java's `Lock` variables are similar to the locks of C#, except that in Java, a lock must be explicitly unlocked with a call to `unlock`. This provides one more way to create erroneous code. C# threads, like those of Java, are lightweight, so although they are more efficient, they cannot

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8. Bags are unordered collections of objects.
be as versatile as Ada’s tasks. The availability of the concurrent collection classes is another advantage C# has over the other nonfunctional languages discussed in this chapter.

13.9 Concurrency in Functional Languages

This section provides a brief overview of support for concurrency in several functional programming languages.

13.9.1 Multi-LISP

Multi-LISP (Halstead, 1985) is an extension to Scheme that allows the programmer to specify program parts that can be executed concurrently. These forms of concurrency are implicit; the programmer is simply telling the compiler (or interpreter) some parts of the program that can be run concurrently.

One of the ways a programmer can tell the system about possible concurrency is the \texttt{pcall} construct. If a function call is embedded in a \texttt{pcall} construct, the parameters to the function can be evaluated concurrently. For example, consider the following \texttt{pcall} construct:

\begin{verbatim}
(pcall f a b c d)
\end{verbatim}

The function is \texttt{f}, with parameters \texttt{a}, \texttt{b}, \texttt{c}, and \texttt{d}. The effect of \texttt{pcall} is that the parameters of the function can be evaluated concurrently (any or all of the parameters could be complicated expressions). Unfortunately, whether this process can be safely used, that is, without affecting the semantics of the function evaluation, is the responsibility of the programmer. This is actually a simple matter if the language does not allow side effects or if the programmer designed the function not to have side effects or at least to have limited ones. However, Multi-LISP does allow some side effects. If the function was not written to avoid side effects, it may be difficult for the programmer to determine whether \texttt{pcall} can be safely used.

The \texttt{future} construct of Multi-LISP is a more interesting and potentially more productive source of concurrency. As with \texttt{pcall}, a function call is wrapped in a \texttt{future} construct. Such a function is evaluated in a separate thread, with the parent thread continuing its execution. The parent thread continues until it needs to use the return value of the function. If the function has not completed its execution when its result is needed, the parent thread waits until it has before it continues.

If a function has two or more parameters, they can also be wrapped in \texttt{future} constructs, in which case their evaluations can be done concurrently in separate threads.

These are the only additions to Scheme in Multi-LISP.
### 13.9.2 Concurrent ML

Concurrent ML (CML) is an extension to ML that includes a form of threads and a form of synchronous message passing to support concurrency. The language is completely described in Reppy (1999).

A thread is created in CML with the `spawn` primitive, which takes the function as its parameter. In many cases, the function is specified as an anonymous function. As soon as the thread is created, the function begins its execution in the new thread. The return value of the function is discarded. The effects of the function are either output produced or through communications with other threads. Either the parent thread (the one that spawned the new thread) or the child thread (the new one) could terminate first and it would not affect the execution of the other.

Channels provide the means of communicating between threads. A channel is created with the `channel` constructor. For example, the following statement creates a channel of arbitrary type named `mychannel`:

```ml
let val mychannel = channel()
```

The two primary operations (functions) on channels are for sending (`send`) and receiving (`recv`) messages. The type of the message is inferred from the send operation. For example, the following function call sends the integer value 7, and therefore the type of the channel is then inferred to be integer:

```ml
send(mychannel, 7)
```

The `recv` function names the channel as its parameter. Its return value is the value it received.

Because CML communications are synchronous, a message is both sent and received only if both the sender and the receiver are ready. If a thread sends a message on a channel and no other thread is ready to receive on that channel, the sender is blocked and waits for another thread to execute a `recv` on the channel. Likewise, if a `recv` is executed on a channel by a thread but no other thread has sent a message on that channel, the thread that ran the `recv` is blocked and waits for a message on that channel.

Because channels are types, functions can take them as parameters.

As was the case with Ada’s synchronous message passing, an issue with CML synchronous message passing is deciding which message to choose when more than one channel has received one. And the same solution is used: the guarded command `do-od` construct that chooses randomly among messages to different channels.

The synchronization mechanism of CML is the `event`. An explanation of this complicated mechanism is beyond the scope of this chapter (and this book).
13.9.3 F#

Part of the F# support for concurrency is based on the same .NET classes that are used by C#, specifically System.Threading.Thread. For example, suppose we want to run the function myConMethod in its own thread. The following function, when called, will create the thread and start the execution of the function in the new thread:

```fsharp
let createThread() =
    let newThread = new Thread(myConMethod)
    newThread.Start()
```

Recall that in C#, it is necessary to create an instance of a predefined delegate, ThreadStart, send its constructor the name of the subprogram, and send the new delegate instance as a parameter to the Thread constructor. In F#, if a function expects a delegate as its parameter, a lambda expression or a function can be sent and the compiler will behave as if you sent the delegate. So, in the above code, the function myConMethod is sent as the parameter to the Thread constructor, but what is actually sent is a new instance of ThreadStart (to which was sent myConMethod).

The Thread class defines the Sleep method, which puts the thread from which it is called to sleep for the number of milliseconds that is sent to it as a parameter.

Shared immutable data does not require synchronization among the threads that access it. However, if the shared data is mutable, which is possible in F#, locking will be required to prevent corruption of the shared data by multiple threads attempting to change it. A mutable variable can be locked while a function operates on it to provide synchronized access to the object with the lock function. This function takes two parameters, the first of which is the variable to be changed. The second parameter is a lambda expression that changes the variable.

A mutable heap-allocated variable is of type ref. For example, the following declaration creates such a variable named sum with the initial value of 0:

```fsharp
let sum = ref 0
```

A ref type variable can be changed in a lambda expression that uses the ALGOL/Pascal/Ada assignment operator, :=. The ref variable must be prefixed with an exclamation point (!) to get its value. In the following, the mutable variable sum is locked while the lambda expression adds the value of x to it:

```fsharp
lock(sum) (fun () -> sum := !sum + x)
```

Threads can be called asynchronously, just as with C#, using the same subprograms, BeginInvoke and EndInvoke, as well as the IAsyncResult interface to facilitate the determination of the completion of the execution of the asynchronously called thread.
As stated previously, F# has the concurrent generic collections of .NET available to its programs. This can save a great deal of programming effort when building multithreaded programs that need a shared data structure in the form of a queue, stack, or bag.

### 13.10 Statement-Level Concurrency

In this section, we take a brief look at language design for statement-level concurrency. From the language design point of view, the objective of such designs is to provide a mechanism that the programmer can use to inform the compiler of ways it can map the program onto a multiprocessor architecture.\(^9\)

In this section, only one collection of linguistic constructs from one language for statement-level concurrency is discussed: High-Performance Fortran.

#### 13.10.1 High-Performance Fortran

High-Performance Fortran (HPF; ACM, 1993b) is a collection of extensions to Fortran 90 that are meant to allow programmers to specify information to the compiler to help it optimize the execution of programs on multiprocessor computers. HPF includes both new specification statements and intrinsic, or built-in, subprograms. This section discusses only some of the HPF statements.

The primary specification statements of HPF are for specifying the number of processors, the distribution of data over the memories of those processors, and the alignment of data with other data in terms of memory placement. The HPF specification statements appear as special comments in a Fortran program. Each of them is introduced by the prefix `!HPFS`, where `!` is the character used to begin lines of comments in Fortran 90. This prefix makes them invisible to Fortran 90 compilers but easy for HPF compilers to recognize.

The `PROCESSORS` specification has the following form:

```fortran
!HPFS PROCESSORS procs (n)
```

This statement is used to specify to the compiler the number of processors that can be used by the code generated for this program. This information is used in conjunction with other specifications to tell the compiler how data are to be distributed to the memories associated with the processors.

The `DISTRIBUTE` and `ALIGN` specifications are used to provide information to the compiler on machines that do not share memory—that is, each processor has its own memory. The assumption is that an access by a processor to its own memory is faster than an access to the memory of another processor.

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9. Although ALGOL 68 included a semaphore type that was meant to deal with statement-level concurrency, we do not discuss that application of semaphores here.
The \texttt{DISTRIBUTE} statement specifies what data are to be distributed and the kind of distribution that is to be used. Its form is as follows:

\begin{verbatim}
!HPF$ DISTRIBUTE (kind) ONTO procs :: identifier_list
\end{verbatim}

In this statement, \texttt{kind} can be either \texttt{BLOCK} or \texttt{CYCLIC}. The identifier list is the names of the array variables that are to be distributed. A variable that is specified to be \texttt{BLOCK} distributed is divided into \(n\) equal groups, where each group consists of contiguous collections of array elements evenly distributed over the memories of all the processors. For example, if an array with 500 elements named \texttt{LIST} is \texttt{BLOCK} distributed over 5 processors, the first 100 elements of \texttt{LIST} will be stored in the memory of the first processor, the second 100 in the memory of the second processor, and so forth. A \texttt{CYCLIC} distribution specifies that individual elements of the array are cyclically stored in the memories of the processors. For example, if \texttt{LIST} is \texttt{CYCLIC} distributed, again over five processors, the first element of \texttt{LIST} will be stored in the memory of the first processor, the second element in the memory of the second processor, and so forth.

The form of the \texttt{ALIGN} statement is

\begin{verbatim}
ALIGN array1_element WITH array2_element
\end{verbatim}

\texttt{ALIGN} is used to relate the distribution of one array with that of another. For example,

\begin{verbatim}
ALIGN list1(index) WITH list2(index+1)
\end{verbatim}

specifies that the \texttt{index} element of \texttt{list1} is to be stored in the memory of the same processor as the \texttt{index+1} element of \texttt{list2}, for all values of \texttt{index}. The two array references in an \texttt{ALIGN} appear together in some statement of the program. Putting them in the same memory (which means the same processor) ensures that the references to them will be as close as possible.

Consider the following example code segment:

\begin{verbatim}
REAL list_1 (1000), list_2 (1000)
INTEGER list_3 (500), list_4 (501)
!HPF$ PROCESSORS proc (10)
!HPF$ DISTRIBUTE (BLOCK) ONTO procs :: list_1, list_2
!HPF$ ALIGN list_3 (index) WITH list_4 (index+1)

\ldots

list_1 (index) = list_2 (index)
list_3 (index) = list_4 (index+1)
\end{verbatim}

In each execution of these assignment statements, the two referenced array elements will be stored in the memory of the same processor.

The HPF specification statements provide information for the compiler that it may or may not use to optimize the code it produces. What the compiler actually does depends on its level of sophistication and the particular architecture of the target machine.
The **FORALL** statement specifies a sequence of assignment statements that may be executed concurrently. For example,

```fortran
FORALL (index = 1:1000)
  list_1(index) = list_2(index)
END FORALL
```

specifies the assignment of the elements of `list_2` to the corresponding elements of `list_1`. However, the assignments are restricted to the following order: the right side of all 1,000 assignments must be evaluated first, before any assignments take place. This permits concurrent execution of all of the assignment statements. In addition to assignment statements, **FORALL** statements can appear in the body of a **FORALL** construct. The **FORALL** statement is a good match with vector machines, in which the same instruction is applied to many data values, usually in one or more arrays. The HPF **FORALL** statement is included in Fortran 95 and subsequent versions of Fortran.

We have briefly discussed only a small part of the capabilities of HPF. However, it should be enough to provide the reader with an idea of the kinds of language extensions that are useful for programming computers with possibly large numbers of processors.

**C# 4.0** (and the other .NET languages) include two methods that behave somewhat like **FORALL**. They are loop control statements in which the iterations can be unrolled and the bodies executed concurrently. These are `Parallel.For` and `Parallel.ForEach`.

**Summary**

Concurrent execution can be at the instruction, statement, or subprogram level. We use the phrase *physical concurrency* when multiple processors are actually used to execute concurrent units. If concurrent units are executed on a single processor, we use the term *logical concurrency*. The underlying conceptual model of all concurrency can be referred to as *logical concurrency*.

Most multiprocessor computers fall into one of two broad categories—SIMD or MIMD. MIMD computers can be distributed.

Languages that support subprogram-level concurrency must provide two fundamental capabilities: mutually exclusive access to shared data structures (competition synchronization) and cooperation among tasks (cooperation synchronization).

Tasks can be in any one of five different states: new, ready, running, blocked, or dead.

Rather than designing language constructs for supporting concurrency, sometimes libraries, such as OpenMP, are used.

The design issues for language support for concurrency are how competition and cooperation synchronization are provided, how an application can
influence task scheduling, how and when tasks start and end their executions, and how and when they are created.

A semaphore is a data structure consisting of an integer and a task description queue. Semaphores can be used to provide both competition and cooperation synchronization among concurrent tasks. It is easy to use semaphores incorrectly, resulting in errors that cannot be detected by the compiler, linker, or run-time system.

Monitors are data abstractions that provide a natural way of providing mutually exclusive access to data shared among tasks. They are supported by several programming languages, among them Ada, Java, and C#. Cooperation synchronization in languages with monitors must be provided with some form of semaphores.

The underlying concept of the message-passing model of concurrency is that tasks send each other messages to synchronize their execution.

Ada provides complex but effective constructs, based on the message-passing model, for concurrency. Ada’s tasks are heavyweight tasks. Tasks communicate with each other through the rendezvous mechanism, which is synchronous message passing. A rendezvous is the action of a task accepting a message sent by another task. Ada includes both simple and complicated methods of controlling the occurrences of rendezvous among tasks.

Ada 95+ includes additional capabilities for the support of concurrency, primarily protected objects. Ada 95+ supports monitors in two ways, with tasks and with protected objects.

Java supports lightweight concurrent units in a relatively simple but effective way. Any class that either inherits from Thread or implements Runnable can override a method named run and have that method’s code executed concurrently with other such methods and with the main program. Competition synchronization is specified by defining methods that access shared data to be implicitly synchronized. Small sections of code can also be implicitly synchronized. A class whose methods are all synchronized is a monitor. Cooperation synchronization is implemented with the methods wait, notify, and notifyAll. The Thread class also provides the sleep, yield, join, and interrupt methods.

Java has direct support for counting semaphores through its Semaphore class and its acquire and release methods. It also had some classes for providing nonblocking atomic operations, such as addition, increment, and decrement operations for integers. Java also provides explicit locks with the Lock interface and ReentrantLock class and its lock and unlock methods. In addition to implicit synchronization using synchronized, Java provides implicit nonblocking synchronization of int, long, and boolean type variables, as well as references and arrays. In these cases, atomic getters, setters, add, increment, and decrement operations are provided.

C#’s support for concurrency is based on that of Java but is slightly more sophisticated. Any method can be run in a thread. Both actor and server threads are supported. All threads are controlled through associated delegates. Server threads can be synchronously called with Invoke or asynchronously called
with BeginInvoke. A callback method address can be sent to the called thread. Three kinds of thread synchronization are supported with the Interlocked class, which provides atomic increment and decrement operations, the Monitor class, and the lock statement.

All .NET languages have the use of the generic concurrent data structures for stacks, queues, and bags, for which competition synchronization is implicit.

Multi-LISP extends Scheme slightly to allow the programmer to inform the implementation about program parts that can be executed concurrently. Concurrent ML extends ML to support a form of threads and a form of synchronous message passing among those threads. This message passing is designed with channels. F# programs have access to all of the .NET support classes for concurrency. Data shared among threads that is mutable can have access synchronized.

High-Performance Fortran includes statements for specifying how data is to be distributed over the memory units connected to multiple processors. Also included are statements for specifying collections of statements that can be executed concurrently.

**BIBLIOGRAPHIC NOTES**

The general subject of concurrency is discussed at great length in Andrews and Schneider (1983), Holt et al. (1978), and Ben-Ari (1982).

The monitor concept is developed and its implementation in Concurrent Pascal is described by Brinch Hansen (1977).


**REVIEW QUESTIONS**

1. When is a concurrent program scalable?
2. What are the motivations to use concurrent machines?
3. Who suggested a categorization of computer architectures?
4. How many types of computer architectures are there?
5. What level of program concurrency is best supported by MIMD computers?
6. What is the producer–consumer problem?
7. What is the job of a scheduler?
8. What is a thread of control in a program?
9. Why are coroutines called quasi-concurrent?
10. What is a virtually multithreaded program?
11. What are four reasons for studying language support for concurrency?
12. What is a heavyweight task? What is a lightweight task?
13. Define task, synchronization, competition and cooperation synchronization, liveness, race condition, and deadlock.
14. What kind of tasks do not require any kind of synchronization?
15. Describe the five different states in which a task can be.
16. What is a task descriptor?
17. In the context of language support for concurrency, what is a guard?
18. What is the purpose of a task-ready queue?
19. Mention two programming languages that support concurrency.
20. What is a guard in a semaphore?
21. What is a binary semaphore? What is a counting semaphore?
22. What are the primary problems with using semaphores to provide synchronization?
23. What advantage do monitors have over semaphores?
24. In what three common languages can monitors be implemented?
25. Define rendezvous, accept clause, entry clause, actor task, server task, extended accept clause, open accept clause, closed accept clause, and completed task.
26. Which is more general, concurrency through monitors or concurrency through message passing?
27. What are the two operations used in semaphores?
28. What purpose does an extended accept clause serve?
29. How is cooperation synchronization provided for Ada tasks?
30. What is the advantage of protected objects in Ada 95 over tasks for providing access to shared data objects?
31. Specifically, what Java program unit can run concurrently with the main method in an application program?
32. Mention any two programming languages that support monitors.
33. What predefined class is designed to implement monitors?
34. What does the Java yield method do?
35. What does the Java join method do?
36. What does the Java interrupt method do?
37. What are the two Java constructs that can be declared to be synchronized?
38. What are the maximum and minimum priorities of a thread in Java?
39. Can Java threads be actor threads, server threads, or either?
40. Describe the actions of the three Java methods that are used to support cooperation synchronization.
41. What kind of Java object is a monitor?
42. Explain why Java includes the Runnable interface.
43. What are the two methods used with Java Semaphore objects?
44. What is the advantage of the nonblocking synchronization in Java?
45. What are the methods of the Java AtomicInteger class and what is the purpose of this class?
46. How are explicit locks supported in Java?
47. Which object is used to create a C# thread?
48. Can C# threads be actor threads, server threads, or either?
49. What are the two ways a C# thread can be called synchronously?
50. How can a C# thread be called asynchronously?
51. How is the returned value from an asynchronously called thread retrieved in C#?
52. What is different about C#’s Sleep method, relative to Java’s sleep?
53. What exactly does C#’s Abort method do?
54. What is the purpose of C#’s Interlocked class?
55. What does the C# lock statement do?
56. On what language is Multi-LISP based?
57. What is the semantics of Multi-LISP’s pcall construct?
58. What is concurrent ML (CML)?
59. What is the type of an F# heap-allocated mutable variable?
60. Why don’t F# immutable variables require synchronized access in a multithreaded program?
61. What is the objective of the specification statements of HPF?
62. What is the purpose of the FORALL statement of HPF and Fortran?

**Problem Set**

1. Explain clearly why competition synchronization is not a problem in a programming environment that supports coroutines but not concurrency.
2. How is deadlock related to liveness?
3. Busy waiting is a method whereby a task waits for a given event by continuously checking for that event to occur. What is the main problem with this approach?
4. In the producer-consumer example of Section 13.3, suppose that we incorrectly replaced the `release(access)` in the consumer process with `wait(access)`. What would be the result of this error on execution of the system?

5. From a book on assembly language programming for a computer that uses an Intel Pentium processor, determine what instructions are provided to support the construction of semaphores.

6. Suppose two tasks, A and B, must use the shared variable Buf_Size. Task A adds 2 to Buf_Size, and task B subtracts 1 from it. Assume that such arithmetic operations are done by the three-step process of fetching the current value, performing the arithmetic, and putting the new value back. In the absence of competition synchronization, what sequences of events are possible and what values result from these operations? Assume that the initial value of Buf_Size is 6.

7. Compare the Java competition synchronization mechanism with that of Ada.

8. Compare the Java cooperation synchronization mechanism with that of Ada.

9. What will be the effect if the wait and signal operations are made nonatomic in semaphores?

10. Explain why the programming languages used for developing operating systems must have threads.

**Programming Exercises**

1. Write an Ada task to implement general semaphores.

2. Write an Ada task to manage a shared buffer such as the one in our example, but use the semaphore task from Programming Exercise 1.

3. Define semaphores in Ada and use them to provide both cooperation and competition synchronization in the shared-buffer example.

4. Write Programming Exercise 3 using Java.

5. Write the shared-buffer example of the chapter in C#.

6. The reader-writer problem can be stated as follows: A shared memory location can be concurrently read by any number of tasks, but when a task must write to the shared memory location, it must have exclusive access. Write a Java program for the reader-writer problem.


8. Write Programming Exercise 6 using C#.
14

Exception Handling and Event Handling

14.1 Introduction to Exception Handling
14.2 Exception Handling in C++
14.3 Exception Handling in Java
14.4 Exception Handling in Python and Ruby
14.5 Introduction to Event Handling
14.6 Event Handling with Java
14.7 Event Handling in C#
This chapter discusses programming language support for two related parts of many contemporary programs: exception handling and event handling. Both exceptions and events can occur at times that cannot be predetermined, and both are best handled with special language constructs and processes. Some of these constructs and processes—for example, propagation—are similar for exception handling and event handling.

We first describe the fundamental concepts of exception handling, including hardware- and software-detectable exceptions, exception handlers, and the raising of exceptions. Then, the design issues for exception handling are introduced and discussed, including the binding of exceptions to exception handlers, continuation, and default handlers. This section is followed by descriptions and evaluations of the exception-handling facilities of two programming languages: C++ and Java. Brief introductions to exception handling in Python and Ruby are then presented.

The latter part of this chapter is about event handling. We first present an introduction to the basic concepts of event handling. This is followed by discussions of the event-handling approaches of Java and C#.

### 14.1 Introduction to Exception Handling

Most computer hardware systems are capable of detecting certain run-time error conditions, such as floating-point overflow. Early programming languages were designed and implemented in such a way that the user program could neither detect nor attempt to deal with such errors. In these languages, the occurrence of such an error simply causes the program to be terminated and control to be transferred to the operating system. The typical operating system reaction to a run-time error is to display a diagnostic message, which may be meaningful and therefore useful, or highly cryptic. After displaying the message, the program is terminated.

In the case of input and output operations, however, the situation is somewhat different. For example, a Fortran `Read` statement can intercept input errors and end-of-file conditions, both of which are detected by the input device hardware. In both cases, the `Read` statement can specify the label of some statement in the user program that deals with the condition. In the case of the end-of-file, the condition obviously is not always considered an error. In most cases, it is nothing more than a signal that one kind of processing is completed and another kind must begin. In spite of the obvious difference between end-of-file and events that are always errors, such as a failed input process, Fortran handles both situations with the same mechanism. Consider the following Fortran `Read` statement:

```fortran
Read(Unit=5, Fmt=1000, Err=100, End=999) Weight
```

The `Err` clause specifies that control is to be transferred to the statement labeled 100 if an error occurs in the read operation. The `End` clause specifies that control is to be transferred to the statement labeled 999 if the read
operation encounters the end of the file. So, Fortran uses simple branches for both input errors and end-of-file.

There is a category of serious errors that are not detectable by hardware but can be detected by code generated by the compiler. For example, array subscript range errors are almost never detected by hardware,\(^1\) but they lead to serious errors that often are not noticed until later in the program execution.

Detection of subscript range errors is sometimes required by the language design. For example, the Java language specification requires Java compilers to generate code to check the correctness of every subscript expression (they do not generate such code when it can be determined at compile time that a subscript expression cannot have an out-of-range value, for example, if the subscript is a literal). In C, subscript ranges are not checked because the cost of such checking was (and still is) not believed to be worth the benefit of detecting such errors. In some compilers for some languages, subscript range checking can be selected (if not turned on by default) or turned off (if it is on by default) as desired in the program or in the command that executes the compiler.

The designers of most contemporary languages have included mechanisms that allow programs to react in a standard way to certain runtime errors, as well as other program-detected unusual events. Programs may also be notified when certain events are detected by hardware or system software, so that they also can react to these events.

One reason some languages do not include exception handling is the complexity it adds to the language.

### 14.1.1 Basic Concepts

We consider both the errors detected by hardware, such as disk read errors, and unusual conditions, such as end-of-file (which is also detected by hardware), to be exceptions. We further extend the concept of an exception to include errors or unusual conditions that are software-detectable (by either a software interpreter or the user code itself). Accordingly, we define exception to be any unusual event, erroneous or not, that is detectable by either hardware or software and that may require special processing.

The special processing that may be required when an exception is detected is called exception handling. This processing is done by a code unit or segment called an exception handler. An exception is raised when its associated event occurs. In some C-based languages, exceptions are said to be thrown, rather than raised.\(^2\) Different kinds of exceptions require different exception handlers. Detection of end-of-file nearly always requires some specific program action. But, clearly, that action would not also be appropriate for an array index range error exception. In some cases, the only action is the generation of an error message and an orderly termination of the program.

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1. In the 1970s, there were some computers that did detect subscript range errors in hardware.
2. C++ was the first C-based language that included exception handling. The word throw was used, rather than raise, because the standard C library includes a function named raise.
The absence of separate or specific exception-handling facilities in a language does not preclude the handling of user-defined, software-detected exceptions. Such an exception detected within a program unit is often handled by the unit’s caller. One possible design is to send an auxiliary parameter, which is used as a status variable. The status variable is assigned a value in the called subprogram according to the correctness and/or normalness of the results of its computations. Immediately upon return from the called unit, the caller tests the status variable. If the value indicates that an exception has occurred, the handler, which may reside in the calling unit, can be enacted. Many of the C standard library functions use a variant of this approach: The return values are used as error indicators.

Another possibility is to pass a label parameter to the subprogram. Of course, this approach is possible only in languages that allow labels to be used as parameters. Passing a label allows the called unit to return to a different point in the caller if an exception has occurred. As in the first alternative, the handler is often a segment of the calling unit’s code. This is a common use of label parameters in Fortran.

A third possibility is to have the handler defined as a separate subprogram whose name is passed as a parameter to the called unit. In this case, the handler subprogram is provided by the caller, but the called unit calls the handler when an exception is raised. One problem with this approach is that one is required to send a handler subprogram with every call to every subprogram that takes a handler subprogram as a parameter, whether it is needed or not. Furthermore, to deal with several different kinds of exceptions, several different handler routines would need to be passed, complicating the code.

If it is desirable to handle an exception in the unit in which it is detected, the handler is included as a segment of code in that unit.

There are some definite advantages to having exception handling built into a language. First, without exception handling, the code required to detect error conditions can considerably clutter a program. For example, suppose a subprogram includes expressions that contain 10 references to elements of a matrix named mat, and any one of them could have an index out-of-range error. Further suppose that the language does not require index range checking. Without built-in index range checking, every one of these operations may need to be preceded by code to detect a possible index range error. For example, consider the following reference to an element of mat, which has 10 rows and 20 columns:

```java
if (row >= 0 && row < 10 && col >= 0 && col < 20)
    sum += mat[row][col];
else
    System.out.println("Index range error on mat, row = " +
                        row + " col = " + col);
```

The presence of exception handling in the language would permit the compiler to insert machine code for such checks before every array element access, greatly shortening and simplifying the source program.
Another advantage of language support for exception handling results from exception propagation. Exception propagation allows an exception raised in one program unit to be handled in some other unit in its dynamic or static ancestry. This allows a single exception handler to be used for any number of different program units. This reuse can result in significant savings in development cost, program size, and program complexity.

A language that supports exception handling encourages its users to consider all of the events that could occur during program execution and how they can be handled. This approach is far better than not considering such possibilities and simply hoping nothing will go wrong.

Finally, there are programs in which dealing with nonerroneous but unusual situations can be simplified with exception handling, and in which program structure can become overly convoluted without it.

### 14.1.2 Design Issues

We now explore some of the design issues for an exception-handling system when it is part of a programming language. Such a system might allow both predefined and user-defined exceptions and exception handlers. Note that predefined exceptions are implicitly raised, whereas user-defined exceptions must be explicitly raised by user code. Consider the following skeletal subprogram that includes an exception-handling mechanism for an implicitly raised exception:

```c
void example() {
    . . .
    average = sum / total;
    . . .
    return;
    /* Exception handlers */
    when zero_divide {
        average = 0;
        printf("Error–divisor (total) is zero\n");
    }
    . . .
}
```

The exception of division by zero, which is implicitly raised, causes control to transfer to the appropriate handler, which is then executed.

The first design issue for exception handling is how an exception occurrence is bound to an exception handler. This issue occurs on two different levels. On the unit level, there is the question of how the same exception being raised at different points in a unit can be bound to different handlers within the unit. For example, in the example subprogram, there is a handler for a division-by-zero exception that appears to be written to deal with an occurrence of division by zero in a particular statement (the one shown). But suppose the function includes several other expressions with division operators. For
those operators, this handler would probably not be appropriate. So, it should be possible to bind the exceptions that can be raised by particular statements to particular handlers, even though the same exception can be raised by many different statements.

At a higher level, the binding question arises when there is no exception handler local to the unit in which the exception is raised. In this case, the language designer must decide whether to propagate the exception to some other unit and, if so, where. How this propagation takes place and how far it goes have an important impact on the writability of exception handlers. For example, if handlers must be local, then many handlers must be written, which complicates both the writing and reading of the program. On the other hand, if exceptions are propagated, a single handler might handle the same exception raised in several program units, which may require the handler to be more general than one would prefer.

An issue that is related to the binding of an exception to an exception handler is whether information about the exception is made available to the handler.

After an exception handler executes, either control can transfer to somewhere in the program outside of the handler code or program execution can simply terminate. We term this the question of control continuation after handler execution, or simply continuation. Termination is obviously the simplest choice, and in many error exception conditions, the best. However, in other situations, particularly those associated with unusual but not erroneous events, the choice of continuing execution is best. This design is called resumption. In these cases, some conventions must be chosen to determine where execution should continue. It might be the statement that raised the exception, the statement after the statement that raised the exception, or possibly some other unit. The choice to return to the statement that raised the exception may seem like a good one, but in the case of an error exception, it is useful only if the handler somehow is able to modify the values or operations that caused the exception to be raised. Otherwise, the exception will simply be reraised. The required modification for an error exception is often very difficult to predict. Even when possible, however, it may not be a sound practice. It allows the program to remove the symptom of a problem without removing the cause.

The two issues of binding of exceptions to handlers and continuation are illustrated in Figure 14.1.

When exception handling is included, a subprogram’s execution can terminate in two ways: when its execution is complete or when it encounters an exception.3 In some situations, it is necessary to complete some computation

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3. Of course, even if the language does not support exception handling, a subprogram could terminate due to a system-detected error.
regardless of how subprogram execution terminates. The ability to specify such a computation is called **finalization**. The choice of whether to support finalization is obviously a design issue for exception handling.

Another design issue is the following: If users are allowed to define exceptions, how are these exceptions specified? The usual answer is to require that they be declared in the specification parts of the program units in which they can be raised. The scope of a declared exception is usually the scope of the program unit that contains the declaration.

In the case where a language provides predefined exceptions, several other design issues follow. For example, should the language run-time system provide default handlers for the built-in exceptions, or should the user be required to write handlers for all exceptions? Another question is whether predefined exceptions can be raised explicitly by the user program. This usage can be convenient if there are software-detectable situations in which the user would like to use a predefined handler.

Another issue is whether hardware-detectable errors can be handled by user programs. If not, all exceptions obviously are software detectable. A related question is whether there should be any predefined exceptions. Predefined exceptions are implicitly raised by either hardware or system software.

The exception-handling design issues can be summarized as follows:

- How and where are exception handlers specified, and what is their scope?
- How is an exception occurrence bound to an exception handler?
- Can information about an exception be passed to the handler?
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• Where does execution continue, if at all, after an exception handler completes its execution? (This is the question of continuation or resumption.)
• Is some form of finalization provided?
• How are user-defined exceptions specified?
• If there are predefined exceptions, should there be default exception handlers for programs that do not provide their own?
• Can predefined exceptions be explicitly raised?
• Are hardware-detectable errors treated as exceptions that may be handled?
• Are there any predefined exceptions?

We are now in a position to examine the exception-handling facilities of several contemporary programming languages.

14.2 Exception Handling in C++

The exception handling of C++ was accepted by the ANSI C++ standardization committee in 1990 and subsequently found its way into C++ implementations. The design is based in part on the exception handling of CLU, Ada, and ML. In C++, exceptions are user or library defined and explicitly raised.

14.2.1 Exception Handlers

C++ uses a special construct that is introduced with the reserved word `try` to specify the scope for exception handlers. A `try` construct includes a compound statement called the `try clause` and a list of exception handlers. The compound statement defines the scope of the following handlers. The general form of this construct is as follows:

```cpp
try {
    //** Code that might raise an exception
} catch (formal parameter) {
    //** A handler body
}
...
catch (formal parameter) {
    //** A handler body
}
```

Each `catch` function is an exception handler. A `catch` function can have only a single formal parameter, which is similar to a formal parameter in a function definition in C++, including the possibility of it being an ellipsis (`. . .`). A handler with an ellipsis formal parameter is the catch-all handler;
it is enacted for any raised exception if no appropriate handler was found. The formal parameter also can be a naked type specifier, such as `float`, as in a function prototype. In such a case, the only purpose of the formal parameter is to make the handler uniquely identifiable. When information about the exception is to be passed to the handler, the formal parameter includes a variable name that is used for that purpose. Because the class of the parameter can be any user-defined class, the parameter can include as many data members as are necessary. Binding exceptions to handlers are discussed in Section 14.3.2.

In C++, exception handlers can include any C++ code.

### 14.2.2 Binding Exceptions to Handlers

C++ exceptions are raised only by the explicit statement `throw`, whose general form in EBNF is as follows:

```c++
throw [expression];
```

The brackets here are metasymbols used to specify that the expression is optional. A `throw` without an operand can appear only in a handler. When it appears there, it reraises the exception, which is then handled elsewhere.

The type of the `throw` expression selects the particular handler, which of course must have a “matching” type formal parameter. In this case, matching means the following: A handler with a formal parameter of type `T`, `const T`, `T&` (a reference to an object of type `T`), or `const T&` matches a `throw` with an expression of type `T`. In the case where `T` is a class, a handler whose parameter is type `T` or any class that is an ancestor of `T` matches. There are more complicated situations in which a `throw` expression matches a formal parameter, but they are not described here.

An exception raised in a `try` clause causes an immediate end to the execution of the code in that `try` clause. The search for a matching handler begins with the handlers that immediately follow the `try` clause. The matching process is done sequentially on the handlers until a match is found. This means that if any other match precedes an exactly matching handler, the exactly matching handler will not be used. Therefore, handlers for specific exceptions are placed at the top of the list, followed by more generic handlers. The last handler is often one with an ellipsis (`...`) formal parameter, which matches any exception. This would guarantee that all exceptions are caught.

If an exception is raised in a `try` clause and there is no matching handler associated with that `try` clause, the exception is propagated. If the `try` clause is nested inside another `try` clause, the exception is propagated to the handlers associated with the outer `try` clause. If none of the enclosing `try` clauses yields a matching handler, the exception is propagated to the caller of the function in which it was raised. If the call to the function was not in a `try` clause, the exception is propagated to that function’s caller. If no matching handler is found in the program through this propagation process, the default handler is called. This handler is further discussed in Section 14.2.4.
14.2.3 Continuation

After a handler has completed its execution, control flows to the first statement following the try construct (the statement immediately after the last handler in the sequence of handlers of which it is an element). A handler may reraise an exception, using a throw without an expression, in which case that exception is propagated.

14.2.4 Other Design Choices

In terms of the design issues summarized in Section 14.1.2, the exception handling of C++ is simple. There are only user-defined exceptions, and they are not specified (though they might be declared as new classes). There is a default exception handler, unexpected, whose only action is to terminate the program. This handler catches all exceptions not caught by the program. It can be replaced by a user-defined handler. The replacement handler must be a function that returns void and takes no parameters. The replacement function is set by assigning its name to set_terminate.

A C++ function can list the types of the exceptions (the types of the throw expressions) that it could raise. This is done by attaching the reserved word throw, followed by a parenthesized list of these types, to the function header. For example,

```cpp
int fun() throw (int, char *) { . . . }
```

specifies that the function fun could raise exceptions of type int and char * but no others. The purpose of the throw clause is to notify users of the function what exceptions might be raised by the function. The throw clause is in effect a contract between the function and its callers. It guarantees that no other exception will be raised in the function. If the function does throw some unlisted exception, the program will be terminated. Note that the compiler ignores throw clauses.

If the types in the throw clause are classes, then the function can raise any exception that is derived from the listed classes. If a function header has a throw clause and raises an exception that is not listed in the throw clause and is not derived from a class listed there, the default handler is called. Note that this error cannot be detected at compile time. The list of types in the list may be empty, meaning that the function will not raise any exceptions. If there is no throw specification on the header, the function can raise any exception. The list is not part of the function’s type.

If a function overrides a function that has a throw clause, the overriding function cannot have a throw clause with more exceptions than the overridden function.

Although C++ has no predefined exceptions, the standard libraries define and throw exceptions, such as out_of_range, which can be thrown by library container classes, and overflow_error, which can be thrown by math library functions.
14.2.5 An Example

The following example program illustrates some simple uses of exception handlers in C++. The program computes and prints a distribution of input grades by using an array of counters. The input is a sequence of grades, terminated by a negative number. The negative number raises a `NegativeInputException` exception because the grades must be nonnegative integers. There are 10 categories of grades (0 – 9, 10 – 19, . . . , 90 – 100). The grades themselves are used to compute indexes into an array of counters, one for each grade category. Invalid input grades are detected by trapping indexing errors in the counter array. A grade of 100 is special in the computation of the grade distribution because the categories all have 10 possible grade values, except the highest, which has 11 (90, 91, . . . , 100). (The fact that there are more possible A grades than B’s or C’s is conclusive evidence of the generosity of teachers.) The grade of 100 is also handled in the same exception handler that is used for invalid input data.

It produces a distribution of input grades by using an array of counters for 10 categories. Illegal grades are detected by checking for invalid subscripts used in incrementing the selected counter.

```cpp
#include <iostream>
int main() {   //* Any exception can be raised
    int new_grade,
        index,
        limit_1,
        limit_2,
        freq[10] = {0,0,0,0,0,0,0,0,0,0};
    // The exception definition to deal with the end of data
    class NegativeInputException {
    public:
        NegativeInputException() {  //* Constructor
            cout << "End of input data reached" << endl;
        }  //** end of constructor
    }  //** end of NegativeInputException class
    try {
        while (true) {
            cout << "Please input a grade" << endl;
            if ((cin >> new_grade) < 0) //* End of data
                throw NegativeInputException();
            index = new_grade / 10;
            //try {
            if (index > 9)
                throw new_grade;
```
freq[index]++;  
}  //* end of inner try compound

catch(int grade) {  //* Handler for index errors
    if (grade == 100)
        freq[9]++;
    else
        cout << "Error -- new grade: " << grade
        << " is out of range" << endl;
}  //* end of catch(int grade)
}  //*  end of the block for the inner try-catch pair
}  //* end of while (1)
}  //* end of outer try block

catch(NegativeInputException& e) {  //**Handler for
    //** negative input
    cout << "Limits   Frequency" << endl;
    for (index = 0; index < 10; index++) {
        limit_1 = 10 * index;
        limit_2 = limit_1 + 9;
        if (index == 9)
            limit_2 = 100;
        cout << limit_1 << limit_2 << freq[index] << endl;
    }  //* end of for (index = 0)
}  //* end of catch (NegativeInputException& e)
}  //* end of main

This program is meant to illustrate the mechanics of C++ exception handling. Note that the index range exception is often handled in C++ by overloading the indexing operation, which could then raise the exception, rather than the direct detection of the indexing operation with the selection construct used in our example.

### 14.2.6 Evaluation

One deficiency of exception handling in C++ is that there are no predefined hardware-detectable exceptions that can be handled by the user. Exceptions are connected to handlers through a parameter type in which the formal parameter may be omitted. The type of the formal parameter of a handler determines the condition under which it is called but may have nothing whatsoever to do with the nature of the raised exception. Therefore, the use of predefined types for exceptions certainly does not promote readability. It is much better to define classes for exceptions with meaningful names in a meaningful hierarchy that can be used for defining exceptions. The exception parameter provides a way to pass information about an exception to the exception handler.

### 14.3 Exception Handling in Java

In Chapter 13, the Java example program includes the use of exception handling with little explanation. This section describes the details of Java's exception-handling capabilities.
14.3 Exception Handling in Java

Java’s exception handling is based on that of C++, but it is designed to be more in line with the object-oriented language paradigm. Furthermore, Java includes a collection of predefined exceptions that are implicitly raised by the Java run-time system.

14.3.1 Classes of Exceptions

All Java exceptions are objects of classes that are descendants of the Throwable class. The Java system includes two predefined exception classes that are subclasses of Throwable: Error and Exception. The Error class and its descendants are related to errors that are thrown by the Java run-time system, such as running out of heap memory. These exceptions are never thrown by user programs, and they should never be handled there. There are two system-defined direct descendants of Exception: RuntimeException and IOException. As its name indicates, IOException is thrown when an error has occurred in an input or output operation, all of which are defined as methods in the various classes defined in the package java.io.

There are predefined classes that are descendants of RuntimeException. In most cases, RuntimeException is thrown when a user program causes an error. For example, ArrayIndexOutOfBoundsException, which is defined in java.util, is a commonly thrown exception that descends from RuntimeException. Another commonly thrown exception that descends from RuntimeException is NullPointerException.

User programs can define their own exception classes. The convention in Java is that user-defined exceptions are subclasses of Exception.

14.3.2 Exception Handlers

The exception handlers of Java have the same form as those of C++, except that every catch must have a parameter and the class of the parameter must be a descendant of the predefined class Throwable.

The syntax of the try construct in Java is exactly as that of C++, except for the finally clause described in Section 14.3.6.

14.3.3 Binding Exceptions to Handlers

Throwing an exception is simple. An instance of the exception class is given as the operand of the throw statement. For example, suppose we define an exception named MyException as

```java
class MyException extends Exception {
    public MyException() {}
    public MyException(String message) {
        super (message);
    }
}
```
This exception can be thrown with the following statement:

```java
throw new MyException();
```

One of the two constructors we have included in our new class has no parameter and the other has a `String` object parameter that it sends to the superclass (`Exception`), which displays it. Therefore, our new exception could be thrown with

```java
throw new MyException("a message to specify the location of the error");
```

The binding of exceptions to handlers in Java is similar to that of C++. If an exception is thrown in the compound statement of a `try` construct, it is bound to the first handler (`catch` function) immediately following the `try` clause whose parameter is the same class as the thrown object, or an ancestor of it. If a matching handler is found, the `throw` is bound to it and it is executed.

Exceptions can be handled and then rethrown by including a `throw` statement without an operand at the end of the handler. The newly thrown exception will not be handled in the same `try` where it was originally thrown, so looping is not a concern. This rethrowing is usually done when some local action is useful, but further handling by an enclosing `try` clause or a `try` clause in the caller is necessary. A `throw` statement in a handler could also throw some exception other than the one that transferred control to this handler.

To ensure that exceptions that can be thrown in a `try` clause are always handled in a method, a special handler can be written that matches all exceptions that are derived from `Exception` simply by defining the handler with an `Exception` type parameter, as in

```java
catch (Exception genericObject) {
    . . .
}
```

Because a class name always matches itself or any ancestor class, any class derived from `Exception` matches `Exception`. Of course, such an exception handler should always be placed at the end of the list of handlers, for it will block the use of any handler that follows it in the `try` construct in which it appears. This occurs because the search for a matching handler is sequential, and the search ends when a match is found.

### 14.3.4 Other Design Choices

As part of its reflection facilities, the Java run-time system stores the class name of every object in the program. The method `getClass` can be used to get an object that stores the class name, which itself can be gotten with the `getName` method. So, we can retrieve the name of the class of the actual parameter from
the `throw` statement that caused the handler’s execution. For the handler shown earlier, this is done with

```java
genericObject.getClass().getName()
```

In addition, the message associated with the parameter object, which is created by the constructor, can be gotten with

```java
genericObject.getMessage()
```

Furthermore, in the case of user-defined exceptions, the thrown object could include any number of data fields that might be useful in the handler.

The `throws` clause of Java has the appearance and placement (in a program) that is similar to that of the `throw` specification of C++. However, the semantics of `throws` is somewhat different from that of the C++ `throw` clause.

The appearance of an exception class name in the `throws` clause of a Java method specifies that that exception class or any of its descendant exception classes can be thrown but not handled by the method. For example, when a method specifies that it can throw `IOException`, it means it can throw an `IOException` object or an object of any of its descendant classes, such as `EOFException`, and it does not handle the exception it throws.

Exceptions of class `Error` and `RuntimeException` and their descendants are called **unchecked exceptions**. All other exceptions are called **checked exceptions**. Unchecked exceptions are never a concern of the compiler. However, the compiler ensures that all checked exceptions a method can throw are either listed in its `throws` clause or handled in the method. Note that checking this at compile time differs from C++, in which it is done at run time. The reason why exceptions of the classes `Error` and `RuntimeException` and their descendants are unchecked is that any method could throw them. A program can catch unchecked exceptions, but it is not required.

As is the case with C++, a method cannot declare more exceptions in its `throws` clause than the method it overrides, though it may declare fewer. So if a method has no `throws` clause, neither can any method that overrides it. A method can throw any exception listed in its `throws` clause, along with any of the descendant classes of those exceptions.

A method that does not directly throw a particular exception, but calls another method that could throw that exception, must list the exception in its `throws` clause. This is the reason the `buildDist` method (in the example in the next subsection), which uses the `readLine` method, must specify `IOException` in the `throws` clause of its header.

A method that does not include a `throws` clause cannot propagate any checked exception. Recall that in C++, a function without a `throw` clause can throw any exception.

A method that calls a method that lists a particular checked exception in its `throws` clause has three alternatives for dealing with that exception: First, it can catch the exception and handle it. Second, it can catch the exception and throw an exception that is listed in its own `throws` clause. Third, it could
declare the exception in its own `throws` clause and not handle it, which effectively propagates the exception to an enclosing `try` clause, if there is one, or to the method’s caller, if there is no enclosing `try` clause.

There are no default exception handlers, and it is not possible to disable exceptions. Continuation in Java is exactly as in C++.

### 14.3.5 An Example

Following is the Java program with the capabilities of the C++ program in Section 14.2.5:

```java
// Grade Distribution
// Input: A list of integer values that represent grades, followed by a negative number
// Output: A distribution of grades, as a percentage for each of the categories 0-9, 10-19, . . . , 90-100.
import java.io.*;
// The exception definition to deal with the end of data
class NegativeInputException extends Exception {
    public NegativeInputException() {
        System.out.println("End of input data reached");
    }  //** end of constructor
}  //** end of NegativeInputException class

class GradeDist {
    int newGrade,
        index,
        limit_1,
        limit_2;
    int [] freq = {0, 0, 0, 0, 0, 0, 0, 0, 0, 0};

    void buildDist() throws IOException {
        DataInputStream in = new DataInputStream(System.in);
        try {
            while (true) {
                System.out.println("Please input a grade");
                newGrade = Integer.parseInt(in.readLine());
                if (newGrade < 0)
                    throw new NegativeInputException();
                index = newGrade / 10;
                try {
                    freq[index]++;
                }  //** end of inner try clause
                catch (ArrayIndexOutOfBoundsException e) {
                    if (newGrade == 100)
                        freq [9]++;
                    else
                        System.out.println("Error - new grade: " + newGrade + " is out of range");
                }
            }
        }
```
The exception for a negative input, `NegativeInputException`, is defined in the program. Its constructor displays a message when an object of the class is created. Its handler produces the output of the method. `ArrayIndexOutOfBoundsException` is a predefined unchecked exception that is thrown by the Java run-time system. In both of these cases, the handler does not include an object name in its parameter. In neither case would a name serve any purpose. Although all handlers get objects as parameters, they often are not useful.

14.3.6 The `finally` Clause

There are some situations in which a process must be executed regardless of whether a `try` clause throws an exception or the exception is handled in the method. One example of such a situation is a file that must be closed. Another is if the method has some external resource that must be freed in the method regardless of how the execution of the method terminates. The `finally` clause was designed for these kinds of needs. A `finally` clause is placed at the end of the list of handlers just after a complete `try` construct. In general, the `try` construct and its `finally` clause appear as

```java
try {
    . . .
}
catch ( . . . ) {
    . . .
} /// More handlers
finally {
    . . .
}
```

The semantics of this construct is as follows: If the `try` clause throws no exceptions, the `finally` clause is executed before execution continues after the `try` construct. If the `try` clause throws an exception and it is caught by a following handler, the
finally clause is executed after the handler completes its execution. If the try clause throws an exception but it is not caught by a handler following the try construct, the finally clause is executed before the exception is propagated.

A try construct with no exception handlers can be followed by a finally clause. This makes sense, of course, only if the compound statement has a throw, break, continue, or return statement. Its purpose in these cases is the same as when it is used with exception handling. For example, consider the following:

```java
try {
    for (index = 0; index < 100; index++) {
        . . .
        if (. . . ) {
            return;
        }  //** end of if
        . . .
    }  //** end of for
}  //** end of try clause
finally {
    . . .
}  //** end of try construct
```

The finally clause here will be executed, regardless of whether the return terminates the loop or it ends normally.

### 14.3.7 Assertions

In the discussion of Plankalkül in Chapter 2, we mentioned that it included assertions. Assertions were added to Java in version 1.4. To use them, it is necessary to enable them by running the program with the enableassertions (or ea) flag, as in

```
java -enableassertions MyProgram
```

There are two possible forms of the assert statement:

- `assert condition;`
- `assert condition : expression;`

In the first case, the condition is tested when execution reaches the assert. If the condition evaluates to true, nothing happens. If it evaluates to false, the AssertionError exception is thrown. In the second case, the action is the same, except that the value of the expression is passed to the AssertionError constructor as a string and becomes debugging output.

The assert statement is used for defensive programming. A program may be written with many assert statements, which ensure that the program's computation is on track to produce correct results. Many programmers put in such checks when they write a program, as an aid to debugging, even though the language they are using does not support assertions. When the program
is sufficiently tested, these checks are removed. The advantage of `assert` statements, which have the same purpose, is that they can be disabled without removing them from the program. This saves the effort of removing them and also allows their use during subsequent program maintenance.

### 14.3.8 Evaluation

The Java mechanisms for exception handling are an improvement over the C++ version on which they are based.

First, a C++ program can throw any type defined in the program or by the system. In Java, only objects that are instances of `Throwable` or some class that descends from it can be thrown. This separates the objects that can be thrown from all of the other objects (and nonobjects) that inhabit a program. What significance can be attached to an exception that causes an `int` value to be thrown?

Second, a C++ program unit that does not include a `throw` clause can throw any exception, which tells the reader nothing. A Java method that does not include a `throws` clause cannot throw any checked exception that it does not handle. Therefore, the reader of a Java method knows from its header what exceptions it could throw but does not handle. A C++ compiler ignores `throw` clauses, but a Java compiler ensures that all exceptions that a method can throw are listed in its `throws` clause.

Third, the `finally` clause is a useful addition. It allows cleanup kinds of actions to take place regardless of how a compound statement terminated.

Finally, the Java run-time system implicitly throws a variety of predefined exceptions, such as for array indices out of range and dereferencing null references, which can be handled by any user program. A C++ program can handle only those exceptions that it explicitly throws (or that are thrown by library classes it uses).

C# includes exception-handling constructs that are very much like those of Java, except that C# does not have a `throws` clause.

### 14.4 Exception Handling in Python and Ruby

This section provides brief overviews of the exception-handling mechanisms of Python and Ruby.

#### 14.4.1 Python

In Python, exceptions are objects. The base class of all exception classes is `BaseException`, from which the `Exception` class is derived. `BaseException` provides some services that are useful for all exception classes, but it is not usually directly subclassed. All predefined exception classes are derived from `Exception` and user-defined exception classes also are derived from it. The most commonly used predefined subclasses of `Exception` are `ArithmeticError`,
whose primary subclasses are OverflowError, ZeroDivisionError, and FloatingPointError, and LookupError, whose main subclasses are IndexError and KeyError.

The statements for dealing with exceptions are similar to those of Java. The general form of a try construct is as follows:

```python
try:
    The try block (the range of statements to be watched for exceptions)
except Exception1:
    Handler for Exception1
except Exception2:
    Handler for Exception2
...
else:
    The else block (what to do when no exception is raised)
finally:
    The finally block (what must be done regardless of what happened)
```

Both the else and the finally clauses are optional.

One difference between handlers in Java and Python is that Python uses except to introduce them, rather than catch. The else clause is executed if no exception is raised in the try block. The finally clause has the same semantics as its counterpart in Java: If an exception is raised in the try block but is not handled by an immediately following handler, the exception is propagated after the finally block is executed. Because a handler handles its named exception, as well as all subclasses of that exception, a handler that names Exception handles all predefined and user-defined exceptions.

An unhandled exception is propagated to progressively larger enclosing try constructs, searching for an appropriate handler. If none is found, the exception is propagated to the function’s caller, again searching for a handler in a nesting try construct. If no handler is found at any level, the default handler is called, which produces an error message and a stack trace and terminates the program.

The raise statement of Python is similar to the throw statement of Java and C++. The parameter for raise is the class name of the exception to be raised. For example, we could have the following:

```python
raise IndexError
```

This statement implicitly creates an instance of the named class, IndexError.

An exception handler can gain access to the object of the raised exception by providing an as clause and a variable name, as in the following:

```python
except Exception as ex_obj:
```

This is a universal handler, as it handles all exceptions. The exception object can be printed with a print statement in the handler, which produces the
message of the object. For example, if the exception was `ZeroDivisionError`, the message would be `division by zero`.

Python’s `assert` statement provides a mechanism for making some exception handling optional. The general form of `assert` is as follows:

```python
assert test, data
```

In this statement, `test` is a Boolean flag or expression and `data` is the value that is sent to the constructor for the exception object to be raised. The meaning of this statement, which optionally raises the `AssertionError` exception, can be described with the following code:

```python
if __debug__:
    if not test:
        raise AssertionError(data)
```

`__debug__` is a predefined flag that is set to `True` unless the `-0` flag is used on the command that runs the program. This allows one to disable all `assert` statements for a particular run of the program. If an `AssertionError` exception is not handled by the program, like other unhandled exceptions, it terminates the program after using the default handler.

Python does not have an equivalent to the `throws` clause of Java.

### 14.4.2 Ruby

Like Python, Ruby exceptions are objects and it has a large collection of predefined exception classes. All of the exceptions that are handled by application programs are either objects of the `StandardError` class or a class that descends from it. `StandardError` is derived from `Exception`, which provides two useful methods to all its descendants. These are `message`, which returns the human-readable error message, and `backtrace`, which returns a stack trace starting from the method where the exception was raised. Some of the predefined subclasses of `StandardError` are `ArgumentError`, `IndexError`, `IOError`, and `ZeroDivisionError`.

Exceptions are explicitly raised with the `raise` method. `raise` is often called with a string parameter. In this case, it raises a new `RuntimeError` object with the string as its message. For example, we could have the following:

```ruby
raise "bad parameter" if count == 0
```

`raise` could also have two parameters, the first of which would be an object of an exception class. The `exception` method of this object is called and the returned `Exception` object is raised. In this case, the second parameter would be the string message to be displayed. For example, we could have the following:

```ruby
raise TypeError, "Float parameter expected" if not param.is_a? Float
```
An exception handler is specified with a `rescue` clause, which is attached to a statement. To attach an exception handler to a segment of code, the code is placed in a `begin-end` block. The `rescue` clause is placed in the block after the code of the block. In general, this appears as in the following:

```ruby
begin
  The sequence of statements in the block
rescue
  The handler
end
```

A `begin-end` block can include an `else` clause and/or an `ensure` clause. The `else` clause is exactly like that of Python. The `ensure` clause is exactly like a `finally` clause. A method can act as a container for exception handling in place of a `begin-end` block.

In a clear departure from most other languages, Ruby allows a segment of code that raised an exception to be rerun after the exception is handled. This is specified with a `retry` statement at the end of the handler.

### 14.5 Introduction to Event Handling

Event handling is similar to exception handling. In both cases, the handlers are implicitly called by the occurrence of something, either an exception or an event. While exceptions can be raised either explicitly by user code or implicitly by hardware or a software interpreter, events are created by external actions, such as user interactions through a graphical user interface (GUI). In this section, the fundamentals of event handling, which are less complex than those of exception handling, are introduced.

In conventional (non-event-driven) programming, the program code itself specifies the order in which that code is executed, although the order is usually affected by the program’s input data. In event-driven programming, parts of the program are executed at completely unpredictable times, often triggered by user interactions with the executing program.

The particular kind of event handling discussed in this chapter is related to GUls. Therefore, most of the events are caused by user interactions through graphical objects or components, often called widgets. The most common widgets are buttons. Implementing reactions to user interactions with GUI components is the most common form of event handling.

An `event` is a notification that something specific has occurred, such as a mouse click on a graphical button. Strictly speaking, an event is an object that is implicitly created by the run-time system in response to a user action, at least in the context in which event handling is being discussed here.

An `event handler` is a segment of code that is executed in response to the appearance of an event. Event handlers enable a program to be responsive to user actions.
Although event-driven programming was being used long before GUIs appeared, it has become a widely used programming methodology only in response to the popularity of these interfaces. As an example, consider the GUIs presented to users of Web browsers. Many Web documents presented to browser users are now dynamic. Such a document may present an order form to the user, who chooses the merchandise by clicking buttons. The required internal computations associated with these button clicks are performed by event handlers that react to the click events.

Another common use of event handlers is to check for simple errors and omissions in the elements of a form, either when they are changed or when the form is submitted to the Web server for processing. Using event handling on the browser to check the validity of form data saves the time of sending that data to the server, where their correctness then must be checked by a server-resident program or script before they can be processed. This kind of event-driven programming is often done using a client-side scripting language, such as JavaScript.

### 14.6 Event Handling with Java

In addition to Web applications, non-Web Java applications can present GUIs to users. GUIs in Java applications are discussed in this section.

The initial version of Java provided a somewhat primitive form of support for GUI components. In version 1.2 of the language, released in late 1998, a new collection of components was added. These were collectively called Swing.

#### 14.6.1 Java Swing GUI Components

The Swing collection of classes and interfaces, defined in `javax.swing`, includes GUI components, or widgets. Because our interest here is event handling, not GUI components, we discuss only two kinds of widgets: text boxes and radio buttons.

A text box is an object of class `JTextField`. The simplest `JTextField` constructor takes a single parameter, the length of the box in characters. For example,

```java
JTextField name = new JTextField(32);
```

The `JTextField` constructor can also take a literal string as an optional first parameter. This string parameter, when present, is displayed as the initial contents of the text box.

Radio buttons are special buttons that are placed in a button group container. A button group is an object of class `ButtonGroup`, whose constructor takes no parameters. In a radio button group, only one button can be pressed at a time. If any button in the group becomes pressed, the previously pressed button is implicitly unpressed. The `JRadioButton` constructor, used for creating radio

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4. Over the last few years, Swing is slowly being replaced by a new GUI toolset named JAVAFX.
buttons, takes two parameters: a label and the initial state of the radio button (true or false, for pressed and not pressed, respectively). If one radio button in a group is initially set to pressed, the others in the group default to unpressed. After the radio buttons are created, they are placed in their button group with the add method of the group object. Consider the following example:

```java
ButtonGroup payment = new ButtonGroup();
JRadioButton box1 = new JRadioButton("Visa", true);
JRadioButton box2 = new JRadioButton("Master Charge");
JRadioButton box3 = new JRadioButton("Discover");
payment.add(box1);
payment.add(box2);
payment.add(box3);
```

A JFrame object is a frame, which is displayed as a separate window. The JFrame class defines the data and methods that are needed for frames. So, a class that uses a frame can be a subclass of JFrame. A JFrame has several layers called panes. We are interested in just one of those layers, the content pane. Components of a GUI are placed in a JPanel object (a panel), which is used to organize and define the layout of the components. A frame is created and the panel containing the components is added to that frame’s content pane.

Predefined graphic objects, such as GUI components, are placed directly in a panel. The following creates the panel object used in the following discussion of components:

```java
JPanel myPanel = new JPanel();
```

After the components have been created with constructors, they are placed in the panel with the add method, as in

```java
myPanel.add(button1);
```

### 14.6.2 The Java Event Model

When a user interacts with a GUI component, for example by clicking a button, the component creates an event object and calls an event handler through an object called an event listener, passing the event object. The event handler provides the associated actions. GUI components are event generators. In Java, events are connected to event handlers through event listeners. Event listeners are connected to event generators through event listener registration. Listener registration is done with a method of the class that implements the listener interface, as described later in this section. Only event listeners that are registered for a specific event are notified when that event occurs.

The listener method that receives the message implements an event handler. To make the event-handling methods conform to a standard protocol, an interface is used. An interface prescribes standard method protocols but does not provide implementations of those methods.
A class that needs to implement an event handler must implement an interface for the listener for that handler. There are several classes of events and listener interfaces. One class of events is ItemEvent, which is associated with the event of clicking a checkbox or a radio button, or selecting a list item. The ItemListener interface includes the protocol of a method, itemStateChanged, which is the handler for ItemEvent events. So, to provide an action that is triggered by a radio button click, the interface ItemListener must be implemented, which requires a definition of the method, itemStateChanged.

As stated previously, the connection of a component to an event listener is made with a method of the class that implements the listener interface. For example, because ItemEvent is the class name of event objects created by user actions on radio buttons, the addItemListener method is used to register a listener for radio buttons. The listener for button events created in a panel could be implemented in the panel or a subclass of JPanel. So, for a radio button named button1 in a panel named myPanel that implements the ItemEvent event handler for buttons, we would register the listener with the following statement:

```java
button1.addItemListener(this);
```

Each event handler method receives an event parameter, which provides information about the event. Event classes have methods to access that information. For example, when called through a radio button, the isSelected method returns true or false, depending on whether the button was on or off (pressed or not pressed), respectively.

All the event-related classes are in the java.awt.event package, so it is imported to any class that uses events.

The following is an example application, RadioB, that illustrates the use of events and event handling. This application constructs radio buttons that control the font style of the contents of a text field. It creates a Font object for each of four font styles. Each of these has a radio button to enable the user to select the font style.

The purpose of this example is to show how events raised by GUI components can be handled to change the output display of the program dynamically. Because of our narrow focus on event handling, some parts of this program are not explained here.

```java
/* RadioB.java
   An example to illustrate event handling with interactive radio buttons that control the font style of a textfield */

package radiob;
import java.awt.*;
import java.awt.event.*;
import javax.swing.*;

public class RadioB extends JPanel implements ItemListener {
    private JTextField text;
```
private Font plainFont, boldFont, italicFont, boldItalicFont;
private JRadioButton plain, bold, italic, boldItalic;
private ButtonGroup radioButtons;

// The constructor method is where the display is initially built
public RadioB() {

// Create the test text string and set its font
text = new JTextField("In what font style should I appear?", 25);
text.setFont(plainFont);

// Create radio buttons for the fonts and add them to a new button group
plain = new JRadioButton("Plain", true);
bold = new JRadioButton("Bold");
italic = new JRadioButton("Italic");
boldItalic = new JRadioButton("Bold Italic");
radioButtons = new ButtonGroup();
radioButtons.add(plain);
radioButtons.add(bold);
radioButtons.add(italic);
radioButtons.add(boldItalic);

// Create a panel and put the text and the radio buttons in it; then add the panel to the frame
JPanel radioPanel = new JPanel();
radioPanel.add(text);
radioPanel.add(plain);
radioPanel.add(bold);
radioPanel.add(italic);
radioPanel.add(boldItalic);
add(radioPanel, BorderLayout.LINE_START);

// Register the event handlers
plain.addItemListener(this);
bold.addItemListener(this);
italic.addItemListener(this);
boldItalic.addItemListener(this);

// Create the fonts
plainFont = new Font("Serif", Font.PLAIN, 16);
boldFont = new Font("Serif", Font.BOLD, 16);
italicFont = new Font("Serif", Font.ITALIC, 16);
boldItalicFont = new Font("Serif", Font.BOLD + Font.ITALIC, 16);
} // End of the constructor for RadioB
// The event handler
public void itemStateChanged (ItemEvent e) {

// Determine which button is on and set the font accordingly
if (plain.isSelected())
text.setFont(plainFont);
else if (bold.isSelected())
text.setFont(boldFont);
else if (italic.isSelected())
text.setFont(italicFont);
else if (boldItalic.isSelected())
text.setFont(boldItalicFont);
} // End of itemStateChanged

// The main method
public static void main(String[] args) {
// Create the window frame
JFrame myFrame = new JFrame(" Radio button example");

// Create the content pane and set it to the frame
JComponent myContentPane = new RadioB();
myContentPane.setOpaque(true);
myFrame.setContentPane(myContentPane);

// Display the window.
myFrame.pack();
myFrame.setVisible(true);
}
} // End of RadioB

The RadioB.java application produces the screen shown in Figure 14.2.

14.7 Event Handling in C#

Event handling in C# (and in the other .NET languages) is similar to that of Java. .NET provides two approaches to creating GUIs in applications, the original Windows Forms and the more recent Windows Presentation Foundation. The latter is the more sophisticated and complex of the two. Because
our interest is just in event handling, we will use the simpler Windows Forms to discuss our subject.

Using Windows Forms, a C# application that constructs a GUI is created by subclassing the Form predefined class, which is defined in the System.Windows.Forms namespace. This class implicitly provides a window to contain our components. There is no need to build frames or panels explicitly.

Text can be placed in a Label object and radio buttons are objects of the RadioButton class. The size of a Label object is not explicitly specified in the constructor; rather it can be specified by setting the AutoSize data member of the Label object to true, which sets the size according to what is placed in it.

Components can be placed at a particular location in the window by assigning a new Point object to the Location property of the component. The Point class is defined in the System.Drawing namespace. The Point constructor takes two parameters, which are the coordinates of the object in pixels. For example, Point(100, 200) is a position that is 100 pixels from the left edge of the window and 200 pixels from the top. The label of a component is set by assigning a string literal to the Text property of the component. After creating a component, it is added to the form window by sending it to the Add method of the Controls subclass of the form. Therefore, the following code creates a radio button with the label Plain at the (100, 300) position in the output window:

```csharp
private RadioButton plain = new RadioButton();
plain.Location = new Point(100, 300);
plain.Text = "Plain";
Controls.Add(plain);
```

All C# event handlers have the same protocol: the return type is void and the two parameters are of types object and EventArgs. Neither of the parameters needs to be used for a simple situation. An event handler method can have any name. A radio button is tested to determine whether it is clicked with the Boolean Checked property of the button. Consider the following skeletal example of an event handler:

```csharp
private void rb_CheckedChanged (object o, EventArgs e){
    if (plain.Checked) . . .
    . . .
}
```

To register an event, a new EventHandler object must be created. The constructor for this class is sent the name of the handler method. The new object is added to the predefined delegate for the event on the component object (using the += assignment operator). For example, when a radio button changes from unchecked to checked, the CheckedChanged event is raised and the handlers registered on the associated delegate, which is referenced by the name of the event, are called. If the event handler is named rb_CheckedChanged, the following statement would register the handler for the CheckedChanged event on the radio button plain:
Following is the RadioB example from Section 14.6 rewritten in C#. Once again, because our focus is on event handling, we do not explain all of the details of the program.

// RadioB.cs
// An example to illustrate event handling with
// interactive radio buttons that control the font
// style of a string of text

namespace RadioB {

    using System;
    using System.Drawing;
    using System.Windows.Forms;

    public class RadioB : Form {

        private Label text = new Label();
        private RadioButton plain = new RadioButton();
        private RadioButton bold = new RadioButton();
        private RadioButton italic = new RadioButton();
        private RadioButton boldItalic = new RadioButton();

        // Constructor for RadioB
        public RadioB() {

            // Initialize the attributes of the text and radio
            // buttons
            text.AutoSize = true;
            text.Text = "In what font style should I appear?";
            plain.Location = new Point(220, 0);
            plain.Text = "Plain";
            plain.Checked = true;
            bold.Location = new Point(350, 0);
            bold.Text = "Bold";
            italic.Location = new Point(480, 0);
            italic.Text = "Italics";
            boldItalic.Location = new Point(610, 0);
            boldItalic.Text = "Bold/Italics";

            // Add the text and the radio buttons to the form
            Controls.Add(text);
            Controls.Add(plain);
            Controls.Add(bold);
            Controls.Add(italic);
            Controls.Add(boldItalic);

            // Register the event handler for the radio buttons
            plain.CheckedChanged +=
                new EventHandler(rb_CheckedChanged);

        }

    }

}
bold.CheckedChanged +=
    new EventHandler(rb_CheckedChanged);
italic.CheckedChanged +=
    new EventHandler(rb_CheckedChanged);
boldItalic.CheckedChanged +=
    new EventHandler(rb_CheckedChanged);
}

// The main method is where execution begins
static void Main()
{
    Application.EnableVisualStyles();
    Application.SetCompatibleTextRenderingDefault (false);
    Application.Run(new RadioB());
}

// The event handler
private void rb_CheckedChanged (object o,
    EventArgs e) {

    // Determine which button is on and set the font
    // accordingly
    if (plain.Checked)
        text.Font =
            new Font( text.Font.Name, text.Font.Size,
                FontStyle.Regular);
    if (bold.Checked)
        text.Font =
            new Font(text.Font.Name, text.Font.Size,
                FontStyle.Bold);
    if (italic.Checked)
        text.Font =
            new Font(text.Font.Name, text.Font.Size,
                FontStyle.Italic);
    if (boldItalic.Checked)
        text.Font =
            new Font(text.Font.Name, text.Font.Size,
                FontStyle.Italic ^ FontStyle.Bold);
    }

The output from this program is exactly like that shown in Figure 14.2.

**S U M M A RY**

C++ includes no predefined exceptions (except those defined in the standard library). C++ exceptions are objects of a primitive type, a predefined class, or a user-defined class. Exceptions are bound to handlers by connecting the type of
the expression in the `throw` statement to that of the formal parameter of the handler. Handlers all have the same name—`catch`. The C++ `throw` clause of a method lists the types of exceptions that the method could throw.

Java exceptions are objects whose ancestry must trace back to a class that descends from the `Throwable` class. There are two categories of exceptions—checked and unchecked. Checked exceptions are a concern for the user program and the compiler. Unchecked exceptions can occur anywhere and are often ignored by user programs.

The Java `throws` clause of a method lists the checked exceptions that it could throw and does not handle. It must include exceptions that methods it calls could raise and propagate back to its caller.

The Java `finally` clause provides a mechanism for guaranteeing that some code will be executed regardless of how the execution of a `try` compound terminates.

Java now includes an `assert` statement, which facilitates defensive programming.

Python’s exception handling is similar to that of Java, although it adds the `else` clause to the `try` construct. Also, it uses `except` clauses rather than `catch` clauses to define handlers and `raise` instead of `throw`. Access to the data of an exception object is gained by assigning the object to a variable with an `as` clause. Python’s `assert` statement is a conditional `raise`.

Exception handling in Ruby is similar to that of Python. Every exception class has two methods, `message` and `backtrace`. Exceptions are often raised with a `raise` statement with a single string parameter. This creates a new `RuntimeError` object with the string as its message. The `raise` statement can be made conditional by adding a conditional expression to it. The scope of exception handlers usually is specified with a `begin-end` block. Handlers are defined in `rescue` clauses. A `begin-end` block can include an `else` clause and an `ensure` clause, which is like the `finally` clause of Python and Java.

An event is a notification that something has occurred that requires special processing. Events are often created by user interactions with a program through a graphical user interface. Java event handlers are called through event listeners. An event listener must be registered for an event if it is to be notified when the event occurs. Two of the most commonly used event listeners interfaces are `actionPerformed` and `itemStateChanged`.

Windows Forms is the original approach to building GUI components and handling events in .NET languages. A C# application builds a GUI in this approach by subclassing the `Form` class. All .NET event handlers use the same protocol. Event handlers are registered by creating an `EventHandler` object and assigning it to the predefined delegate associated with the GUI object that can raise the event.

**Bibliographic Notes**

One of the most important papers on exception handling that is not connected with a particular programming language is the work by Goodenough (1975). The problems with the PL/I design for exception handling are covered
in MacLaren (1977). Exception handling in C++ is described by Stroustrup (1997). Exception handling in Java is described by Campione et al. (2001).

**REVIEW QUESTIONS**

1. Define *exception*, *exception handler*, *raising an exception*, *continuation*, *finalization*, and *built-in exception*.
2. How did the early programming languages support the exception-handling mechanism?
3. What are the advantages of having support for exception handling built in to a language?
4. What is the reason that some languages do not include exception handling?
5. Give an example of an error that is not detectable by hardware but can be detected by the code generated by the compiler in Java.
6. What is the use of label parameters in Fortran?
7. How does exception propagation work in Java?
8. How does a `throw` statement without an operand work in C++?
9. How can an exception handler be written in C++ so that it handles any exception?
10. What constraint will a function overriding a function that has a `throw` clause face?
11. Does Java include built-in exceptions?
12. Which library functions in C++ support the `overflow_error` exception?
13. What are the two system-defined direct descendants of the `Throwable` class in Java?
14. What package in Java contains the `ArrayIndexOutOfBoundsException`?
15. How can an exception handler be written in Java so that it handles any exception?
16. What is the difference between C++ exception handlers and Java exception handlers?
17. What is the difference between Java exception handlers and Python exception handlers?
18. How can a method deal with exceptions it receives from another method that lists a particular checked exception in its `throw` clause?
19. In Java, when does a `try` clause only have a `finally` clause and no exception handlers?
20. In Java, what clause should a method include to propagate any checked exception?
22. What advantages do `assert` statements have?
23. What happens when an `assert` statement evaluates to false?
24. What does the `message` method of Ruby’s `StandardError` class do?
25. What exactly happens when a `raise` statement with a string parameter is executed?
26. What does the Python `_debug_` flag do?
27. In what ways are exception handling and event handling related?
29. What is event-driven programming?
30. What is the purpose of the Java `JTextField`?
31. What is the purpose of the Java `ButtonGroup`?
32. What object is often used as the event listener in Java GUI applications?
33. What is the origin of the protocol for an event handler in Java?
34. What method is used to register an event handler in Java?
35. Using .NET’s Windows Forms, what namespace is required to build a GUI for a C# application?
36. How is a component positioned in a form using Windows Forms?
37. What is the protocol of a .NET event handler?
38. What class of object must be created to register a .NET event handler?
39. What role do delegates play in the process of registering event handlers?

**Problem Set**

1. What did the designers of C get in return for not requiring subscript range checking?
2. Write a Fortran `Read` statement that transfers the control to statement 200 if an error occurs and transfers the control to statement 950 if the read operation encounters an end-of-file condition.
3. From textbooks on the PL/I and Ada programming languages, look up the respective sets of built-in exceptions. Do a comparative evaluation of the two, considering both completeness and flexibility.
4. From a textbook on COBOL, determine how exception handling is done in COBOL programs.
5. In languages without exception-handling facilities, it is common to have most subprograms include an “error” parameter, which can be set to some value representing “OK” or some other value representing “error in procedure.” What advantage does a linguistic exception-handling facility like that of Java have over this method?
6. In a language without exception-handling facilities, we could send an error-handling procedure as a parameter to each procedure that can detect errors that must be handled. What disadvantages are there to this method?
7. Compare the methods suggested in Problems 5 and 6. Which do you think is better and why?
8. Write a comparative analysis of the `throw` clause of C++ and the `throws` clause of Java.

9. Consider the following C++ skeletal program:

```cpp
class Big {
    int i;
    float f;
    void fun1() throw int {
        . . .
        try {
            . . .
            throw i;
            . . .
            throw f;
            . . .
        }
        catch (float) { . . . }
        . . .
    }
}

class Small {
    int j;
    float g;
    void fun2() throw float {
        . . .
        try {
            . . .
            try {
                Big.fun1();
                . . .
                throw j;
                . . .
                throw g;
                . . .
            }
            catch (int) { . . . }
            . . .
        }
        catch (float) { . . . }
    }
}
```

In each of the four `throw` statements, where is the exception handled?
Note that `fun1` is called from `fun2` in class `Small`.

10. Write a detailed comparison of the exception-handling capabilities of C++ and those of Java.

11. With the help of a book on ML, write a detailed comparison of the exception-handling capabilities of ML and those of Java.

12. Summarize the arguments in favor of the termination and resumption models of continuation.
1. Suppose you are writing a C++ function that has three alternative approaches for accomplishing its requirements. Write a skeletal version of this function so that if the first alternative raises any exception, the second is tried, and if the second alternative raises any exception, the third is executed. Write the code as if the three methods were procedures named `alt1`, `alt2`, and `alt3`.

2. Write a Java program that inputs a list of integer values in the range of \(-100\) to \(100\) from the keyboard and computes the sum of the squares of the input values. This program must use exception handling to ensure that the input values are in range and are legal integers, to handle the error of the sum of the squares becoming larger than a standard `Integer` variable can store, and to detect end-of-file and use it to cause the output of the result. In the case of overflow of the sum, an error message must be printed and the program terminated.

3. Write a C++ program for the specification of Programming Exercise 2.

4. Revise the Java program of Section 14.3.5 to use `EOFException` to detect the end of the input.

5. Rewrite the Java code of Section 14.3.6 that uses a `finally` clause in C++.
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15.1 Introduction
15.2 Mathematical Functions
15.3 Fundamentals of Functional Programming Languages
15.4 The First Functional Programming Language: Lisp
15.5 An Introduction to Scheme
15.6 Common Lisp
15.7 ML
15.8 Haskell
15.9 F#
15.10 Support for Functional Programming in Primarily Imperative Languages
15.11 A Comparison of Functional and Imperative Languages
This chapter introduces functional programming and some of the programming languages that have been designed for this approach to software development. We begin by reviewing the fundamental ideas of mathematical functions, because functional languages are based on them. Next, the idea of a functional programming language is introduced, followed by a look at the first functional language, Lisp. The next, somewhat lengthy section, is devoted to an introduction to Scheme, including some of its primitive functions, special forms, functional forms, and some examples of simple functions written in Scheme. Next, we provide brief introductions to Common Lisp, ML, Haskell, and F#. Then, we discuss support for functional programming that is included in some imperative languages. The following section describes some of the applications of functional programming languages. Finally, we present a brief comparison of functional and imperative languages.

15.1 Introduction

Most of the earlier chapters of this book have been concerned primarily with the imperative programming languages. The high degree of similarity among the imperative languages arises in part from one of the common bases of their design: the von Neumann architecture, as discussed in Chapter 1. Imperative languages can be thought of collectively as a progression of developments to improve the basic model, which was Fortran I. All have been designed to make efficient use of von Neumann architecture computers. Although the imperative style of programming has been found acceptable by most programmers, its heavy reliance on the underlying architecture is thought by some to be an unnecessary restriction on the alternative approaches to software development.

Other bases for language design exist, some of them oriented more to particular programming paradigms or methodologies than to efficient execution on a particular computer architecture. Thus far, however, only a relatively small minority of programs have been written in nonimperative languages.

The functional programming paradigm, which is based on mathematical functions, is the design basis of the most important nonimperative styles of languages. This style of programming is supported by functional programming languages.

The 1977 ACM Turing Award was given to John Backus for his work in the development of Fortran. Each recipient of this award presents a lecture when the award is formally given, and the lecture is subsequently published in the Communications of the ACM. In his Turing Award lecture, Backus (1978) made a case that purely functional programming languages are better than imperative languages because they result in programs that are more readable, more reliable, and more likely to be correct. The crux of his argument was that purely functional programs are easier to understand, both during and after development, largely because the meanings of expressions are independent of their context (one characterizing feature of a pure functional programming language is that neither expressions nor functions have side effects).

In this lecture, Backus proposed a pure functional language, FP (functional programming), which he used to frame his argument. Although the language
did not succeed, at least in terms of achieving widespread use, his idea motivated debate and research on pure functional programming languages. The point here is that some well-known computer scientists have attempted to promote the concept that functional programming languages are superior to the traditional imperative languages, though those efforts have obviously fallen short of their goals. However, over the last decade, prompted in part by the maturing of the typed functional languages, such as ML, Haskell, OCaml, and F#, there has been an increase in the interest in and use of functional programming languages.

One of the fundamental characteristics of programs written in imperative languages is that they have state, which changes throughout the execution process. This state is represented by the program’s variables. The author and all readers of the program must understand the uses of its variables and how the program’s state changes through execution. For a large program, this is a daunting task. This is one problem with programs written in an imperative language that is not present in a program written in a pure functional language, for such programs have neither variables nor state.

Lisp began as a pure functional language but soon acquired some important imperative features that increased its execution efficiency. It is still the most important of the functional languages, at least in the sense that it is the only one that has achieved widespread use. It dominates in the areas of knowledge representation, machine learning, intelligent training systems, and the modeling of speech. Common Lisp is an amalgam of several early 1980s dialects of Lisp.

Scheme is a small, static-scoped dialect of Lisp. Scheme has been widely used to teach functional programming. It is also used in some universities to teach introductory programming courses.

The development of the typed functional programming languages, primarily ML, Haskell, OCaml, and F#, has led to a significant expansion of the areas of computing in which functional languages are now used. As these languages have matured, their practical use is growing. They are now being used in areas such as database processing, financial modeling, statistical analysis, and bioinformatics.

One objective of this chapter is to provide an introduction to functional programming using the core of Scheme, intentionally leaving out its imperative features. Sufficient material on Scheme is included to allow the reader to write some simple but interesting programs. It is difficult to acquire an actual feel for functional programming without some actual programming experience, so that is strongly encouraged.

### 15.2 Mathematical Functions

A mathematical function is a mapping of members of one set, called the **domain set**, to another set, called the **range set**. A function definition specifies the domain and range sets, either explicitly or implicitly, along with the mapping. The mapping is described by an expression or, in some cases, by a
table. Functions are often applied to a particular element of the domain set, given as a parameter to the function. Note that the domain set may be the cross product of several sets (reflecting that there can be more than one parameter). A function yields an element of the range set.

One of the fundamental characteristics of mathematical functions is that the evaluation order of their mapping expressions is controlled by recursion and conditional expressions, rather than by the sequencing and iterative repetition that are common to programs written in the imperative programming languages.

Another important characteristic of mathematical functions is that because they have no side effects and cannot depend on any external values, they always map a particular element of the domain to the same element of the range. However, a subprogram in an imperative language may depend on the current values of several nonlocal or global variables. This makes it difficult to determine statically what values the subprogram will produce and what side effects it will have on a particular execution.

In mathematics, there is no such thing as a variable that models a memory location. Local variables in functions in imperative programming languages maintain the state of the function. Computation is accomplished by evaluating expressions in assignment statements that change the state of the program. In mathematics, there is no concept of the state of a function.

A mathematical function maps its parameter(s) to a value (or values), rather than specifying a sequence of operations on values in memory to produce a value.

### 15.2.1 Simple Functions

Function definitions are often written as a function name, followed by a list of parameters in parentheses, followed by the mapping expression. For example,

\[
cube(x) \equiv x \times x \times x,
\]

where \( x \) is a real number.

In this definition, the domain and range sets are the real numbers. The symbol \( \equiv \) is used to mean “is defined as.” The parameter \( x \) can represent any member of the domain set, but it is fixed to represent one specific element during evaluation of the function expression. This is one way the parameters of mathematical functions differ from the variables in imperative languages.

Function applications are specified by pairing the function name with a particular element of the domain set. The range element is obtained by evaluating the function-mapping expression with the domain element substituted for the occurrences of the parameter. Once again, it is important to note that during evaluation, the mapping of a function contains no unbound parameters, where a bound parameter is a name for a particular value. Every occurrence of a parameter is bound to a value from the domain set and is a constant during evaluation. For example, consider the following evaluation of \( \text{cube}(x) \):

\[
\text{cube}(2.0) = 2.0 \times 2.0 \times 2.0 = 8
\]
The parameter \( x \) is bound to 2.0 during the evaluation and there are no unbound parameters. Furthermore, \( x \) is a constant (its value cannot be changed) during the evaluation.

Early theoretical work on functions separated the task of defining a function from that of naming the function. Lambda notation, as devised by Alonzo Church (1941), provides a method for defining nameless functions. A **lambda expression** specifies the parameters and the mapping of a function. The lambda expression is the function itself, which is nameless. For example, consider the following lambda expression:

\[
\lambda(x)x \ast x \ast x
\]

Church defined a formal computation model (a formal system for function definition, function application, and recursion) using lambda expressions. This is called **lambda calculus**. Lambda calculus can be either typed or untyped. Untyped lambda calculus serves as the inspiration for the functional programming languages.

As stated earlier, before evaluation a parameter represents any member of the domain set, but during evaluation it is bound to a particular member. When a lambda expression is evaluated for a given parameter, the expression is said to be applied to that parameter. The mechanics of such an application are the same as for any function evaluation. Application of the example lambda expression is denoted as in the following example:

\[(\lambda(x)x \ast x \ast x)(2)\]

which results in the value 8.

Lambda expressions, like other function definitions, can have more than one parameter.

### 15.2.2 Functional Forms

A **higher-order function**, or **functional form**, is one that either takes one or more functions as parameters or yields a function as its result, or both. One common kind of functional form is **function composition**, which has two functional parameters and yields a function whose value is the first actual parameter function applied to the result of the second. Function composition is written as an expression, using \( \circ \) as an operator, as in

\[b \equiv f \circ g\]

For example, if

\[f(x) \equiv x + 2\]
\[g(x) \equiv 3 \ast x\]

then \( b \) is defined as

\[b(x) \equiv f(g(x)), \text{ or } b(x) \equiv (3 \ast x) + 2\]
Apply-to-all is a functional form that takes a single function as a parameter.\footnote{In programming languages, these are often called \textit{map} functions.} If applied to a list of parameters, apply-to-all applies its functional parameter to each of the values in the list parameter and collects the results in a list or sequence. Apply-to-all is denoted by $\alpha$. Consider the following example:

Let

$$h(x) \equiv x^*x$$

then

$$\alpha(h,(2,3,4)) \text{ yields } (4, 9, 16)$$

There are other functional forms, but these two examples illustrate the basic characteristics of all of them.

\section*{15.3 Fundamentals of Functional Programming Languages}

The objective of the design of a functional programming language is to mimic mathematical functions to the greatest extent possible. This results in an approach to problem solving that is fundamentally different from approaches used with imperative languages. In an imperative language, an expression is evaluated and the result is stored in a memory location, which is represented as a variable in a program. This is the purpose of assignment statements. This necessary attention to memory cells, whose values represent the state of the program, results in a relatively low-level programming methodology.

A program in an assembly language often must also store the results of partial evaluations of expressions. For example, to evaluate

$$(x + y)/(a - b)$$

the value of $(x + y)$ is computed first. That value must then be stored while $(a - b)$ is evaluated. The compiler handles the storage of intermediate results of expression evaluations in high-level languages. The storage of intermediate results is still required, but the details are hidden from the programmer.

A purely functional programming language does not use variables or assignment statements, thus freeing the programmer from concerns related to the memory cells, or state, of the program. Without variables, iterative constructs are not possible, for they are controlled by variables. Repetition must be specified with recursion rather than with iteration. Programs are function definitions and function application specifications, and executions consist of evaluating function applications. Without variables, the execution of a purely functional program has no state in the sense of operational and denotational semantics. The execution of a function always produces the same result when given the same parameters. This feature is called referential transparency. It makes the semantics of purely functional languages far simpler than the semantics of the imperative languages (and the functional languages that include
imperative features). It also makes testing easier, because each function can be tested separately, without any concern for its context.

A functional language provides a set of primitive functions, a set of functional forms to construct complex functions from those primitive functions, a function application operation, and some structure or structures for representing data. These structures are used to represent the parameters and values computed by functions. If a functional language is well designed, it requires only a relatively small number of primitive functions.

As we have seen in earlier chapters, the first functional programming language, Lisp, uses a syntactic form for both data and code that is very different from that of the imperative languages. However, many functional languages designed later use syntax for their code that is similar to that of the imperative languages.

Although there are a few purely functional languages, for example, Haskell, most of the languages that are called functional include some imperative features, for example, mutable variables and constructs that act as assignment statements.

Some concepts and constructs that originated in functional languages, such as lazy evaluation and anonymous subprograms, have now found their way into some languages that are considered imperative.

Although early functional languages were often implemented with interpreters, many programs written in functional programming languages are now compiled.

15.4 The First Functional Programming Language: Lisp

Many functional programming languages have been developed. The oldest and most widely used is Lisp (or one of its descendants), which was developed by John McCarthy at MIT in 1959. Studying functional languages through Lisp is somewhat akin to studying the imperative languages through Fortran: Lisp was the first functional language, but although it has steadily evolved for half a century, it no longer represents the latest design concepts for functional languages. In addition, with the exception of the first version, all Lisp dialects include imperative-language features, such as imperative-style variables, assignment statements, and iteration. (Imperative-style variables are used to name memory cells, whose values can change many times during program execution.) Despite this and their somewhat odd form, the descendants of the original Lisp represent well the fundamental concepts of functional programming and are therefore worthy of study.

15.4.1 Data Types and Structures

There were only two categories of data objects in the original Lisp: atoms and lists. List elements are pairs, where the first part is the data of the element, which is a pointer to either an atom or a nested list. The second part of a pair can be a pointer to an atom, a pointer to another element, or a special value, nil. Elements are linked together in lists with the second parts. Atoms and lists
are not types in the sense that imperative languages have types. In fact, the original Lisp was a typeless language. Atoms are either symbols, in the form of identifiers, or numeric literals.

Recall from Chapter 2, that Lisp originally used lists as its data structure because they were thought to be an essential part of list processing. As it eventually developed, however, Lisp rarely requires the general list operations of insertion and deletion at positions other than the beginning of a list.

Lists are specified in Lisp by delimiting their elements with parentheses. The elements of simple lists are restricted to atoms, as in

\[(A \ B \ C \ D)\]

Nested list structures are also specified by parentheses. For example, the list

\[(A \ (B \ C) \ D \ (E \ (F \ G)))\]

is a list of four elements. The first is the atom \(A\); the second is the sublist \((B \ C)\); the third is the atom \(D\); the fourth is the sublist \((E \ (F \ G))\), which has as its second element the sublist \((F \ G)\).

In a Lisp implementation, a list is usually stored as linked list structure in which each node has two pointers, one to reference the data of the node and the other to form the linked list. A list is referenced by a pointer to its first element.

The internal representations of our two example lists are shown in Figure 15.1. Note that the elements of a list are shown horizontally. The last

![Figure 15.1](image-url)
element of a list has no successor, so its link is nil. Sublists are shown with the same structure.

### 15.4.2 The First Lisp Interpreter

The original intent of Lisp’s design was to have a notation for programs that would be as close to Fortran’s as possible, with additions when necessary. This notation was called M-notation, for meta-notation. There was to be a compiler that would translate programs written in M-notation into semantically equivalent machine code programs for the IBM 704.

Early in the development of Lisp, McCarthy wrote a paper to promote list processing as an approach to general symbolic processing. McCarthy believed that list processing could be used to study computability, that at the time was usually studied using Turing machines, which are based on the imperative model of computation. McCarthy thought that the functional processing of symbolic lists was a more natural model of computation than Turing machines, that operated on symbols written on tapes, which represented state.

One of the common requirements of the study of computation is that one must be able to prove certain computability characteristics of the whole class of whatever model of computation is being used. In the case of the Turing machine model, one can construct a universal Turing machine that can mimic the operations of any other Turing machine. From this concept came the idea of constructing a universal Lisp function that could evaluate any other function in Lisp.

The first requirement for the universal Lisp function was a notation that allowed functions to be expressed in the same way data was expressed. The parenthesized list notation described in Section 15.4.1 had already been adopted for Lisp data, so it was decided to invent conventions for function definitions and function calls that could also be expressed in list notation. Function calls were specified in a prefix list form originally called Cambridge Polish, as in the following:

```
(function_name parameter1 . . . parameter_n)
```

For example, if $+$ is a function that takes two or more numeric parameters, the following two expressions evaluate to 12 and 20, respectively:

```
(+ 5 7)
(+ 3 4 7 6)
```

The lambda notation described in Section 15.2.1 was chosen to specify function definitions. It had to be modified, however, to allow the binding of

---

2. This name first was used in the early development of Lisp. The name was chosen because Lisp lists resemble the prefix notation used by the Polish logician Jan Lukasiewicz, and because Lisp was born at MIT in Cambridge, Massachusetts. Some now prefer to call the notation Cambridge prefix.
functions to names so that functions could be referenced by other functions and by themselves. This name binding was specified by a list consisting of the function name and a list containing the lambda expression, as in

\[(\text{function\_name} \ (\text{LAMBDA} \ (\text{param}_1... \text{param}_n) \ \text{expression}))\]

If you have had no prior exposure to functional programming, it may seem odd to even consider a nameless function. However, nameless functions are sometimes useful in functional programming (as well as in mathematics and imperative programming). For example, consider a function whose action is to produce a function for immediate application to a parameter list. The produced function has no need for a name, for it is applied only at the point of its construction. Such an example is given in Section 15.5.14.

Lisp functions specified in this new notation were called S-expressions, for symbolic expressions. Eventually, all Lisp structures, both data and code, were called S-expressions. An S-expression can be either a list or an atom. We will usually refer to S-expressions simply as expressions.

McCarthy successfully developed a universal function that could evaluate any other function. This function was named \texttt{EVAL} and was itself in the form of an expression. Two of the people in the AI Project, which was developing Lisp, Stephen B. Russell and Daniel J. Edwards, noticed that an implementation of \texttt{EVAL} could serve as a Lisp interpreter, and they promptly constructed such an implementation (McCarthy et al., 1965).

There were several important results of this quick, easy, and unexpected implementation. First, all early Lisp implementations copied \texttt{EVAL} and were therefore interpretive. Second, the definition of M-notation, which was the planned programming notation for Lisp, was never completed or implemented, so S-expressions became Lisp's only notation. The use of the same notation for data and code has important consequences, one of which will be discussed in Section 15.5.14. Third, much of the original language design was effectively frozen, keeping certain odd features in the language, such as the conditional expression form and the use of ( ) for both the empty list and logical false.

Another feature of early Lisp systems that was apparently accidental was the use of dynamic scoping. Functions were evaluated in the environments of their callers. No one at the time knew much about scoping, and there may have been little thought given to the choice. Dynamic scoping was used for most dialects of Lisp before 1975. Contemporary dialects either use static scoping or allow the programmer to choose between static and dynamic scoping.

An interpreter for Lisp can be written in Lisp. Such an interpreter, which is not a large program, describes the operational semantics of Lisp, in Lisp. This is vivid evidence of the semantic simplicity of the language.

---

3. There are also uses of nameless subprograms in imperative programming.
15.5 An Introduction to Scheme

In this section, we describe the core part of Scheme (Dybvig, 2009). We have chosen Scheme because it is relatively simple, it is popular in colleges and universities, and Scheme interpreters are readily available (and free) for a wide variety of computers. The version of Scheme described in this section is Scheme 4. Note that this section covers only a small part of Scheme, and it includes none of Scheme’s imperative features.

15.5.1 Origins of Scheme

The Scheme language, which is a dialect of Lisp, was developed at MIT in the mid-1970s (Sussman and Steele, 1975). It is characterized by its small size, its exclusive use of static scoping, and its treatment of functions as first-class entities. As first-class entities, Scheme functions can be the values of expressions, elements of lists, passed as parameters, and returned from functions. Early versions of Lisp did not provide all of these capabilities.

As an essentially typeless small language with simple syntax and semantics, Scheme is well suited to educational applications, such as courses in functional programming, and also to general introductions to programming.

Most of the Scheme code in the following sections would require only minor modifications to be converted to valid Lisp code.

15.5.2 The Scheme Interpreter

A Scheme interpreter in interactive mode is an infinite read-evaluate-print loop (often abbreviated as REPL). It repeatedly reads an expression typed by the user (in the form of a list), interprets the expression, and displays the resulting value. This form of interpreter is also used by Ruby and Python. Expressions are interpreted by the function `EVAL`. Literals evaluate to themselves. So, if you type a number to the interpreter, it simply displays the number. Expressions that are calls to primitive functions are evaluated in the following way: First, each of the parameter expressions is evaluated, in no particular order. Then, the primitive function is applied to the parameter values, and the resulting value is displayed.

Of course, Scheme programs that are stored in files can be loaded and interpreted.

Comments in Scheme are any text following a semicolon on any line.

15.5.3 Primitive Numeric Functions

Scheme includes primitive functions for the basic arithmetic operations. These are `+`, `-`, `*`, and `/`, for add, subtract, multiply, and divide. `*` and `/` can have zero or more parameters. If `*` is given no parameters, it returns 1; if `/` is given no parameters, it returns 0. `+` adds all of its parameters together. `*` multiplies all its
parameters together. \(/\) and \(−\) can have two or more parameters. In the case of subtraction, all but the first parameter are subtracted from the first. Division is similar to subtraction. Some examples are:

<table>
<thead>
<tr>
<th>Expression</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>42</td>
<td>42</td>
</tr>
<tr>
<td>((* 3 7))</td>
<td>21</td>
</tr>
<tr>
<td>((+ 5 7 8))</td>
<td>20</td>
</tr>
<tr>
<td>((- 5 6))</td>
<td>-1</td>
</tr>
<tr>
<td>((- 15 7 2))</td>
<td>6</td>
</tr>
<tr>
<td>((- 24 (* 4 3)))</td>
<td>12</td>
</tr>
</tbody>
</table>

There are a large number of other numeric functions in Scheme, among them \texttt{MODULO}, \texttt{ROUND}, \texttt{MAX}, \texttt{MIN}, \texttt{LOG}, \texttt{SIN}, and \texttt{SQRT}. \texttt{SQRT} returns the square root of its numeric parameter, if the parameter's value is not negative. If the parameter is negative, \texttt{SQRT} yields a complex number.

In Scheme, note that we use uppercase letters for all reserved words and predefined functions. The official definition of the language specifies that there is no distinction between uppercase and lowercase in these. However, some implementations, for example DrRacket's teaching languages, require lowercase for reserved words and predefined functions.

If a function has a fixed number of parameters, such as \texttt{SQRT}, the number of parameters in the call must match that number. If not, the interpreter will produce an error message.

### 15.5.4 Defining Functions

A Scheme program is a collection of function definitions. Consequently, knowing how to define these functions is a prerequisite to writing the simplest program. In Scheme, a nameless function actually includes the word \texttt{LAMBDA}, and is called a \texttt{lambda expression}. For example,

\[
\text{(LAMBDA (x) (* x x))}
\]

is a nameless function that returns the square of its given numeric parameter. This function can be applied in the same way that named functions are: by placing it in the beginning of a list that contains the actual parameters. For example, the following expression yields 49:

\[
\text{((LAMBDA (x) (* x x)) 7)}
\]

In this expression, \(x\) is called a \texttt{bound variable} within the lambda expression. During the evaluation of this expression, \(x\) is bound to 7. A bound variable never changes in the expression after being bound to an actual parameter value at the time evaluation of the lambda expression begins.
Lambda expressions can have any number of parameters. For example, we could have the following:

\[(\text{LAMBDA } (a \ b \ c \ x) \ (+ \ (* \ a \ x \ x) \ (* \ b \ x) \ c))\]

The Scheme special form function \texttt{DEFINE} serves two fundamental needs of Scheme programming: to bind a name to a value and to bind a name to a lambda expression. The form of \texttt{DEFINE} that binds a name to a value may make it appear that \texttt{DEFINE} can be used to create imperative language-style variables. However, these name bindings create named values, not variables. 

\texttt{DEFINE} is called a special form because it is interpreted (by \texttt{EVAL}) in a different way than the normal primitives like the arithmetic functions, as we shall soon see.

The simplest form of \texttt{DEFINE} is one used to bind a name to the value of an expression. This form is

\[(\text{DEFINE } \text{symbol} \ \text{expression})\]

For example,

\[(\text{DEFINE pi 3.14159})
(\text{DEFINE two_pi (* 2 pi)})\]

If these two expressions have been typed to the Scheme interpreter and then \texttt{pi} is typed, the number 3.14159 will be displayed; when \texttt{two_pi} is typed, 6.28318 will be displayed. In both cases, the displayed numbers may have more digits than are shown here.

This form of \texttt{DEFINE} is analogous to a declaration of a named constant in an imperative language. For example, in Java, the equivalents to the above defined names are as follows:

\[
\text{final float PI = 3.14159;}
\text{final float TWO_PI = 2.0 * PI;}
\]

Names in Scheme can consist of letters, digits, and special characters except parentheses; they are case insensitive and must not begin with a digit.

The second use of the \texttt{DEFINE} function is to bind a lambda expression to a name. In this case, the lambda expression is abbreviated by removing the word \texttt{LAMBDA}. To bind a name to a lambda expression, \texttt{DEFINE} takes two lists as parameters. The first parameter is the prototype of a function call, with the function name followed by the formal parameters, together in a list. The second list contains an expression to which the name is to be bound. The general form of such a \texttt{DEFINE} is\(^4\)

\[
4. \text{Actually, the general form of \texttt{DEFINE} has as its body a list containing a sequence of one or more expressions, although in most cases only one is included. We include only one for simplicity's sake.}
\]

(DEFINE (function_name parameters)
    (expression)
)

Of course, this form of DEFINE is the definition of a named function.

The following example call to DEFINE binds the name square to a functional expression that takes one parameter:

(DEFINE (square number) (* number number))

After the interpreter evaluates this function, it can be used, as in

(square 5)

which displays 25.

To illustrate the difference between primitive functions and the DEFINE special form, consider the following:

(DEFINE x 10)

If DEFINE were a primitive function, EVAL’s first action on this expression would be to evaluate the two parameters of DEFINE. If x were not already bound to a value, this would be an error. Furthermore, if x were already defined, it would also be an error, because this DEFINE would attempt to redefine x, which is illegal. Remember, x is the name of a value; it is not a variable in the imperative sense.

Following is another example of a function. It computes the length of the hypotenuse (the longest side) of a right triangle, given the lengths of the two other sides.

(DEFINE (hypotenuse side1 side2)
    (SQRT(+(* (square side1) (square side2))))
)

Notice that hypotenuse uses square, which was defined previously.

### 15.5.5 Output Functions

Scheme includes a few simple output functions, but when used with the interactive interpreter, most output from Scheme programs is the normal output from the interpreter, displaying the results of applying EVAL to top-level functions.

Note that explicit input and output are not part of the pure functional programming model, because input operations change the program state and output operations have side effects. Neither of these can be part of a pure functional language. Therefore, this chapter does not describe the explicit input or output functions of Scheme.
15.5.6 Numeric Predicate Functions

A predicate function is one that returns a Boolean value (some representation of either true or false). Scheme includes a collection of predicate functions for numeric data. Among them are the following:

<table>
<thead>
<tr>
<th>Function</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>=</td>
<td>Equal</td>
</tr>
<tr>
<td>&lt;&gt;</td>
<td>Not equal</td>
</tr>
<tr>
<td>&gt;</td>
<td>Greater than</td>
</tr>
<tr>
<td>&lt;</td>
<td>Less than</td>
</tr>
<tr>
<td>&gt;=</td>
<td>Greater than or equal to</td>
</tr>
<tr>
<td>&lt;=</td>
<td>Less than or equal to</td>
</tr>
<tr>
<td>EVEN?</td>
<td>Is it an even number?</td>
</tr>
<tr>
<td>ODD?</td>
<td>Is it an odd number?</td>
</tr>
<tr>
<td>ZERO?</td>
<td>Is it zero?</td>
</tr>
</tbody>
</table>

Notice that the names for all predefined predicate functions that have words for names end with question marks. In Scheme, the two Boolean values are #T and #F (or #t and #f), although the Scheme predefined predicate functions return the empty list, (), for false.

When a list is interpreted as a Boolean, any nonempty list evaluates to true; the empty list evaluates to false. This is similar to the interpretation of integers in C as Boolean values; zero evaluates to false and any nonzero value evaluates to true.

In the interest of readability, all of our example predicate functions in this chapter return #F, rather than ()

The NOT function is used to invert the logic of a Boolean expression.

15.5.7 Control Flow

Scheme uses three different constructs for control flow: one similar to the selection construct of the imperative languages and two based on the evaluation control used in mathematical functions.

The Scheme two-way selector function, named IF, has three parameters: a predicate expression, a then expression, and an else expression. A call to IF has the form

(IF predicate then_expression else_expression)

For example,

(DEFINE (factorial n)
  (IF (<= n 1)
Recall that the multiple selection of Scheme, \texttt{COND}, was discussed in Chapter 8. Following is an example of a simple function that uses \texttt{COND}:

\begin{verbatim}
(DEFINE (leap? year)
  (COND
    ((ZERO? (MODULO year 400)) #T)
    ((ZERO? (MODULO year 100)) #F)
    (ELSE (ZERO? (MODULO year 4)))
  )
)
\end{verbatim}

The following subsections contain additional examples of the use of \texttt{COND}. The third Scheme control mechanism is recursion, which is used, as in mathematics, to specify repetition. Most of the example functions in Section 15.5.10 use recursion.

\section*{15.5.8 List Functions}

One of the more common uses of the Lisp-based programming languages is list processing. This subsection introduces the Scheme functions for dealing with lists. Recall that Scheme’s list operations were briefly introduced in Chapter 6. Following is a more detailed discussion of list processing in Scheme.

Scheme programs are interpreted by the function application function, \texttt{EVAL}. When applied to a primitive function, \texttt{EVAL} first evaluates the parameters of the given function. This action is necessary when the actual parameters in a function call are themselves function calls, which is frequently the case. In some calls, however, the parameters are data elements rather than function references. When a parameter is not a function reference, it obviously should not be evaluated. We were not concerned with this earlier, because numeric literals always evaluate to themselves and cannot be mistaken for function names.

Suppose we have a function that has two parameters, an atom and a list, and the purpose of the function is to determine whether the given atom is in the given list. Neither the atom nor the list should be evaluated; they are literal data to be processed. To avoid evaluating a parameter, it is first given as a parameter to the primitive function \texttt{QUOTE}, which simply returns it without change. The following examples illustrate \texttt{QUOTE}:

\begin{verbatim}
(QUOTE A) returns A
(QUOTE (A B C)) returns (A B C)
\end{verbatim}

Calls to \texttt{QUOTE} are usually abbreviated by preceding the expression to be quoted with an apostrophe (‘) and leaving out the parentheses around the expression. Thus, instead of \texttt{(QUOTE (A B))}, ‘(A B) is used.

The necessity of \texttt{QUOTE} arises because of the fundamental nature of Scheme (and the other Lisp-based languages): data and code have the same
form. Although this may seem odd to imperative language programmers, it results in some interesting and powerful processes, one of which is discussed in Section 15.5.14.

The CAR, CDR, and CONS functions were introduced in Chapter 6. Following are additional examples of the operations of CAR and CDR:

(CAR ' (A B C)) returns A
(CAR ' ((A B) C D)) returns (A B)
(CAR ' A) is an error because A is not a list
(CAR ' (A)) returns A
(CAR ' ()) is an error
(CDR ' (A B C)) returns (B C)
(CDR ' ((A B) C D)) returns (C D)
(CDR ' A) is an error
(CDR ' (A)) returns ()
(CDR ' ()) is an error

The names of the CAR and CDR functions are peculiar at best. The origin of these names lies in the first implementation of Lisp, which was on an IBM 704 computer. The 704’s memory words had two fields, named decrement and address, that were used in various operand addressing strategies. Each of these fields could store a machine memory address. The 704 also included two machine instructions, also named CAR (contents of the address part of a register) and CDR (contents of the decrement part of a register), that extracted the associated fields. It was natural to use the two fields to store the two pointers of a list node so that a memory word could neatly store a node. Using these conventions, the CAR and CDR instructions of the 704 provided efficient list selectors. The names carried over into the primitives of all dialects of Lisp.

As another example of a simple function, consider

(DEFINE (second a_list) (CAR (CDR a_list)))

Once this function is evaluated, it can be used, as in

(second ' (A B C))

which returns B.

Some of the most commonly used functional compositions in Scheme are built in as single functions. For example, (CAAR x) is equivalent to (CAR (CAR x)), (CADR x) is equivalent to (CAR (CDR x)), and (CADDAR x) is equivalent to (CAR (CDR (CDR (CAR x)))) . Any combination of A’s and D’s, up to four, are legal between the ‘C’ and the ‘R’ in the function’s name. As an example, consider the following evaluation of CADDAR:

(CADDAR ' ((A B (C) D) E)) =
(CAR (CDR (CDR (CAR ' ((A B (C) D) E))))) =
(CAR (CDR (CDR ' (A B (C) D)))) =
Following are example calls to `CONS`:

- `(CONS 'A () )` returns `(A)`
- `(CONS 'A '(B C))` returns `(A B C)`
- `(CONS '() '(A B))` returns `(() A B)`
- `(CONS '(A B) '(C D))` returns `((A B) C D)`

The results of these `CONS` operations are shown in Figure 15.2. Note that `CONS` is, in a sense, the inverse of `CAR` and `CDR`. `CAR` and `CDR` take a list apart, and `CONS` constructs a new list from two given list parts. The two parameters to `CONS` become the `CAR` and `CDR` of the new list. Thus, if `a_list` is a list, then

```lisp
(CONS (CAR a_list) (CDR a_list))
```

returns a list with the same structure and same elements as `a_list`. 

---

**Figure 15.2**

The result of several `CONS` operations
Dealing only with the relatively simple problems and programs discussed in this chapter, it is unlikely one would intentionally apply CONS to two atoms, although that is legal. The result of such an application is a dotted pair, so named because of the way it is displayed by Scheme. For example, consider the following call:

(CONS 'A 'B)

If the result of this is displayed, it would appear as

(A . B)

This dotted pair indicates that instead of an atom and a pointer or a pointer and a pointer, this cell has two atoms.

LIST is a function that constructs a list from a variable number of parameters. It is a shorthand version of nested CONS functions, as illustrated in the following:

(LIST 'apple 'orange 'grape)

returns

(apple orange grape)

Using CONS, the call to LIST above is written as follows:

(CONS 'apple (CONS 'orange (CONS 'grape '())))

15.5.9 Predicate Functions for Symbolic Atoms and Lists

Scheme has three fundamental predicate functions, EQ?, NULL?, and LIST?, for symbolic atoms and lists.

The EQ? function takes two expressions as parameters, although it is usually used with two symbolic atom parameters. It returns #T if both parameters have the same pointer value—that is, they point to the same atom or list; otherwise, it returns #F. If the two parameters are symbolic atoms, EQ? returns #T if they are the same symbols (because Scheme does not make duplicates of symbols); otherwise #F. Consider the following examples:

(EQ? 'A 'A) returns #T
(EQ? 'A 'B) returns #F
(EQ? 'A ' (A B)) returns #F
(EQ? ' (A B) ' (A B)) returns #F or #T
(EQ? 3.4 (+ 3 0.4)) returns #F or #T

As the fourth example indicates, the result of comparing lists with EQ? is not consistent. The reason for this is that two lists that are exactly the same often are not duplicated in memory. At the time the Scheme system creates a list,
it checks to see whether there is already such a list. If there is, the new list is nothing more than a pointer to the existing list. In these cases, the two lists will be judged equal by \texttt{EQ?}. However, in some cases, it may be difficult to detect the presence of an identical list, in which case a new list is created. In this scenario, \texttt{EQ?} yields \texttt{#F}.

The last case shows that the addition may produce a new value, in which case it would not be equal (with \texttt{EQ?}) to \texttt{3.4}, or it may recognize that it already has the value \texttt{3.4} and use it, in which case \texttt{EQ?} will use the pointer to the old \texttt{3.4} and return \texttt{#T}.

As we have seen, \texttt{EQ?} works for symbolic atoms but does not necessarily work for numeric atoms. The \texttt{=} predicate works for numeric atoms but not symbolic atoms. As discussed previously, \texttt{EQ?} also does not work reliably for list parameters.

Sometimes it is convenient to be able to test two atoms for equality when it is not known whether they are symbolic or numeric. For this purpose, Scheme has a different predicate, \texttt{EQV?}, which works on both numeric and symbolic atoms. Consider the following examples:

\begin{verbatim}
(EQV? 'A 'A) returns #T
(EQV? 'A 'B) returns #F
(EQV? 3 3) returns #T
(EQV? 'A 3) returns #F
(EQV? 3.4 (+ 3 0.4)) returns #T
(EQV? 3.0 3) returns #F
\end{verbatim}

Notice that the last example demonstrates that floating-point values are different from integer values. \texttt{EQV?} is not a pointer comparison, it is a value comparison.

The primary reason to use \texttt{EQ?} or \texttt{=} rather than \texttt{EQV?} when it is possible is that \texttt{EQ?} and \texttt{=} are faster than \texttt{EQV?}.

The \texttt{LIST?} predicate function returns \texttt{#T} if its single argument is a list and \texttt{#F} otherwise, as in the following examples:

\begin{verbatim}
(LIST? '(X Y)) returns #T
(LIST? 'X) returns #F
(LIST? '() ) returns #T
\end{verbatim}

The \texttt{NULL?} function tests its parameter to determine whether it is the empty list and returns \texttt{#T} if it is. Consider the following examples:

\begin{verbatim}
(NULL? '(A B)) returns #F
(NULL? '()) returns #T
(NULL? 'A) returns #F
(NULL? '(())) returns #F
\end{verbatim}

The last call yields \texttt{#F} because the parameter is not the empty list. Rather, it is a list containing a single element, the empty list.
15.5.10 Example Scheme Functions

This section contains several examples of function definitions in Scheme. These programs solve simple list-processing problems.

Consider the problem of membership of a given atom in a given list that does not include sublists. Such a list is called a simple list. If the function is named member, it could be used as follows:

\[
\begin{align*}
\text{(member 'B '(A B C)) returns #T} \\
\text{(member 'B '(A C D E)) returns #F}
\end{align*}
\]

Thinking in terms of iteration, the membership problem is simply to compare the given atom and the individual elements of the given list, one at a time in some order, until either a match is found or there are no more elements in the list to be compared. A similar process can be accomplished using recursion. The function can compare the given atom with the CAR of the list. If they match, the value #T is returned. If they do not match, the CAR of the list should be ignored and the search continued on the CDR of the list. This can be done by having the function call itself with the CDR of the list as the list parameter and return the result of this recursive call. This process will end if the given atom is found in the list. If the atom is not in the list, the function will eventually be called (by itself) with a null list as the actual parameter. That event must force the function to return #F. In this process, there are two ways out of the recursion: Either the list is empty on some call, in which case #F is returned, or a match is found and #T is returned.

Altogether, there are three cases that must be handled in the function: an empty input list, a match between the atom and the CAR of the list, or a mismatch between the atom and the CAR of the list, which causes the recursive call. These three are the three parameters to COND, with the last being the default case that is triggered by an ELSE predicate. The complete function follows:5

\[
\begin{align*}
\text{(DEFINE (member atm a_list)} \\
\text{ (COND)} \\
\text{ \quad ((NULL? a_list) #F)} \\
\text{ \quad ((EQ? atm (CAR a_list)) #T)} \\
\text{ \quad (ELSE (member atm (CDR a_list)))})}
\end{align*}
\]

This form is typical of simple Scheme list-processing functions. In such functions, the data in lists are processed one element at a time. The individual elements are specified with CAR, and the process is continued using recursion on the CDR of the list.

Note that the null test must precede the equal test, because applying CAR to an empty list is an error.

---

5. Most Scheme systems define a function named member and do not allow a user to redefine it. So, if the reader wants to try this function, it must be defined with some other name.
As another example, consider the problem of determining whether two
given lists are equal. If the two lists are simple, the solution is relatively easy,
although some programming techniques with which the reader may not be
familiar are involved. A predicate function, `equalsimp`, for comparing simple
lists is shown here:

```lisp
(define (equalsimp list1 list2)
  (cond
    ((null? list1) (null? list2))
    ((null? list2) #f)
    ((eq? (car list1) (car list2))
     (equalsimp (cdr list1) (cdr list2)))
    (else #f))
)
```

The first case, which is handled by the first parameter to `cond`, is for when
the first list parameter is the empty list. This can occur in an external call if the
first list parameter is initially empty. Because a recursive call uses the `cdr`s of
the two parameter lists as its parameters, the first list parameter can be empty
in such a call (if the first list parameter is now empty). When the first list
parameter is empty, the second list parameter must be checked to see whether
it is also empty. If so, they are equal (either initially or the cars were equal on
all previous recursive calls), and `null?` correctly returns `#t`. If the second list
parameter is not empty, it is larger than the first list parameter and `#f` should
be returned, as it is by `null?`.

The next case deals with the second list being empty when the first list
is not. This situation occurs only when the first list is longer than the second.
Only the second list must be tested, because the first case catches all instances
of the first list being empty.

The third case is the recursive step that tests for equality between two
Corresponding elements in the two lists. It does this by comparing the cars
of the two nonempty lists. If they are equal, then the two lists are equal up to
that point, so recursion is used on the cdrs of both. This case fails when two
unequal atoms are found. When this occurs, the process need not continue, so
the default case `else` is selected, which returns `#f`.

Note that `equalsimp` expects lists as parameters and does not operate
correctly if either or both parameters are atoms.

The problem of comparing general lists is slightly more complex than this,
because sublists must be traced completely in the comparison process. In this
situation, the power of recursion is uniquely appropriate, because the form
of sublists is the same as that of the given lists. Any time the corresponding
elements of the two given lists are lists, they are separated into their two parts,
car and cdr, and recursion is used on them. This is a perfect example of the
usefulness of the divide-and-conquer approach. If the corresponding elements
of the two given lists are atoms, they can simply be compared using `eq?`. 
The definition of the complete function follows:

```
(define (equal list1 list2)
  (cond
    ((not (list? list1)) (eq? list1 list2))
    ((not (list? list2)) #f)
    ((null? list1) (null? list2))
    ((null? list2) #f)
    ((equal (car list1) (car list2))
     (equal (cdr list1) (cdr list2)))
    (else #f))
)
```

The first two cases of the `cond` handle the situation where either of the parameters is an atom instead of a list. The third and fourth cases are for the situation where one or both lists are empty. These cases also prevent subsequent cases from attempting to apply `car` to an empty list. The fifth `cond` case is the most interesting. The predicate is a recursive call with the cars of the lists as parameters. If this recursive call returns `#t`, then recursion is used again on the `cdrs` of the lists. This algorithm allows the two lists to include sublists to any depth.

This definition of `equal` works on any pair of expressions, not just lists. `equal` is equivalent to the system predicate function `equal?`. Note that `equal?` should be used only when necessary (the forms of the actual parameters are not known), because it is much slower than `eq?` and `eqv?`.

Another commonly needed list operation is that of constructing a new list that contains all of the elements of two given list arguments. This is usually implemented as a Scheme function named `append`. The result list can be constructed by repeated use of `cons` to place the elements of the first list argument into the second list argument, which becomes the result list. To clarify the action of `append`, consider the following examples:

```
(append '(A B) '(C D R))  returns  (A B C D R)
(append '(((A B) C) 'D (E F))) returns ((A B) C D (E F))
```

The definition of `append` is 6

```
(define (append list1 list2)
  (cond
    ((null? list1) list2)
    (else (cons (car list1) (append (cdr list1) list2))))
)
```

The first `cond` case is used to terminate the recursive process when the first argument list is empty, returning the second list. In the second case (the `else`),

6. As was the case with `member`, a user usually cannot define a function named `append`.
the **CAR** of the first parameter list is **CONS**ed onto the result returned by the recursive call, which passes the **CDR** of the first list as its first parameter.

Consider the following Scheme function, named **guess**, which uses the **member** function described in this section. Try to determine what it does before reading the description that follows it. Assume the parameters are simple lists.

```
(DEFINE (guess list1 list2)
  (COND
    ((NULL? list1) '())
    ((member (CAR list1) list2)
      (CONS (CAR list1) (guess (CDR list1) list2)))
    (ELSE (guess (CDR list1) list2)))
)
```

**guess** yields a simple list that contains the common elements of its two parameter lists. So, if the parameter lists represent sets, **guess** computes a list that represents the intersection of those two sets.

### 15.5.11 LET

**LET** is a function (initially described in Chapter 5) that creates a local scope in which names are temporarily bound to the values of expressions. It is often used to factor out the common subexpressions from more complicated expressions. These names can then be used in the evaluation of another expression, but they cannot be rebound to new values in **LET**. The following example illustrates the use of **LET**. It computes the roots of a given quadratic equation, assuming the roots are real.\(^7\) The mathematical definitions of the real (as opposed to complex) roots of the quadratic equation \(ax^2 + bx + c\) are as follows: 

\[
\text{root}_1 = \frac{-b + \sqrt{b^2 - 4ac}}{2a} \quad \text{and} \quad \text{root}_2 = \frac{-b - \sqrt{b^2 - 4ac}}{2a}
\]

```
(DEFINE (quadratic_roots a b c)
  (LET (
    (root_part_over_2a
      (/ (SQRT (- (* b b) (* 4 a c))) (* 2 a)))
    (minus_b_over_2a (/ (- 0 b) (* 2 a)))
  )
  (LIST (+ minus_b_over_2a root_part_over_2a)
        (- minus_b_over_2a root_part_over_2a))
)
)
```

This example uses **LIST** to create the list of the two values that make up the result.

Because the names bound in the first part of a **LET** construct cannot be changed in the following expression, they are not the same as local variables in

\(^7\) Some versions of Scheme include “complex” as a data type and will compute the roots of the equation, regardless of whether they are real or complex.
a block in an imperative language. They could all be eliminated by textual substitution of their respective expressions for their names in the \texttt{LET} expression.

\texttt{LET} is actually shorthand for a \texttt{LAMBDA} expression applied to a parameter. The following two expressions are equivalent:

\begin{verbatim}
(LET ((alpha 7)) (* 5 alpha))
((LAMBDA (alpha) (* 5 alpha)) 7)
\end{verbatim}

In the first expression, 7 is bound to \texttt{alpha} with \texttt{LET}; in the second, 7 is bound to \texttt{alpha} through the parameter of the \texttt{LAMBDA} expression.

\subsection{Tail Recursion in Scheme}

A function is \texttt{tail recursive} if its recursive call is the last operation in the function. This means that the return value of the recursive call is the return value of the nonrecursive call to the function. For example, the member function of Section 15.5.10, repeated here, is tail recursive.

\begin{verbatim}
(DEFINE (member atm a_list)
  (COND
   ((NULL? a_list) #F)
   ((EQ? atm (CAR a_list)) #T)
   (ELSE (member atm (CDR a_list)))
  ))
\end{verbatim}

This function can be automatically converted by a compiler to use iteration, resulting in faster execution than in its recursive form.

However, many functions that use recursion for repetition are not tail recursive. Programmers who were concerned with efficiency have discovered ways to rewrite some of these functions so that they are tail recursive. One example of this uses an accumulating parameter and a helper function. As an example of this approach, consider the factorial function from Section 15.5.7, which is repeated here:

\begin{verbatim}
(DEFINE (factorial n)
  (IF (<= n 1)
      1
      (* n (factorial (- n 1))))
)
\end{verbatim}

The last operation of this function is the multiplication. The function works by creating the list of numbers to be multiplied together and then doing the multiplications as the recursion unwinds to produce the result. Each of these numbers is created by an activation of the function and each is stored in an activation record instance. As the recursion unwinds the numbers are multiplied together. Recall that the stack is shown after several recursive calls to factorial in Chapter 9. This factorial function can be rewritten with an auxiliary helper
function, which uses a parameter to accumulate the partial factorial. The helper function, which is tail recursive, also takes factorial’s parameter. These functions are as follows:

```
(DEFINE (facthelper n factpartial)
  (IF (<= n 1)
      factpartial
      (facthelper (- n 1) (* n factpartial)))
)
(DEFINE (factorial n)
  (facthelper n 1)
)
```

With these functions, the result is computed during the recursive calls, rather than as the recursion unwinds. Because there is nothing useful in the activation record instances, they are not necessary. Regardless of how many recursive calls are requested, only one activation record instance is necessary. This makes the tail-recursive version far more efficient than the non-tail-recursive version.

The Scheme language definition requires that Scheme language processing systems convert all tail-recursive functions to replace that recursion with iteration. Therefore, it is important, at least for efficiency’s sake, to define functions that use recursion to specify repetition to be tail recursive. Some optimizing compilers for some functional languages can even perform conversions of some non-tail-recursive functions to equivalent tail-recursive functions and then code these functions to use iteration instead of recursion for repetition.

### 15.5.13 Functional Forms

This section describes two common mathematical functional forms that are provided by Scheme: composition and apply-to-all. Both are mathematically defined in Section 15.2.2.

#### 15.5.13.1 Functional Composition

Functional composition is the only primitive functional form provided by the original Lisp. All subsequent Lisp dialects, including Scheme, also provide it. As stated in Section 15.2.2, function composition is a functional form that takes two functions as parameters and returns a function that first applies the second parameter function to its parameter and then applies the first parameter function to the return value of the second parameter function. In other words, the function \( h \) is the composition function of \( f \) and \( g \) if \( h(x) = f(g(x)) \). For example, consider the following example:

```
(DEFINE (g x) (* 3 x))
(DEFINE (f x) (+ 2 x))
```
Now the functional composition of \( f \) and \( g \) can be written as follows:

\[
\text{(DEFINE (h x) (+ 2 (* 3 x)))}
\]

In Scheme, the functional composition function \textit{compose} can be written as follows:

\[
\text{(DEFINE (compose f g) (LAMBDA (x)(f (g x))))}
\]

For example, we could have the following:

\[
\text{((compose CAR CDR) '((a b) c d))}
\]

This call would yield \( c \). This is an alternative, though less efficient, form of \textit{CADR}. Now consider another call to \textit{compose}:

\[
\text{((compose CDR CAR) '((a b) c d))}
\]

This call would yield \( (b) \). This is an alternative to \textit{CDAR}.

As yet another example of the use of \textit{compose}, consider the following:

\[
\text{(DEFINE (third a_list)}
\text{ ((compose CAR (compose CDR CDR)) a_list))}
\]

This is an alternative to \textit{CADDR}.

15.5.13.2 An Apply-to-All Functional Form

The most common functional forms provided in functional programming languages are variations of mathematical apply-to-all functional forms. The simplest of these is \textit{map}, which has two parameters: a function and a list. \textit{map} applies the given function to each element of the given list and returns a list of the results of these applications. A Scheme definition of \textit{map} follows:

\[
\text{(DEFINE (map fun a_list)}
\text{ (COND}
\text{ ((NULL? a_list) '())}
\text{ (ELSE (CONS (fun (CAR a_list)) (map fun (CDR a_list))))})
\]

Note the simple form of \textit{map}, which expresses a complex functional form.

As an example of the use of \textit{map}, suppose we want all of the elements of a list cubed. We can accomplish this with the following:

\[
\text{(map (LAMBDA (num) (* num num num)) '(3 4 2 6))}
\]

---

8. As was the case with member, \textit{map} is a predefined function that cannot be redefined by users.
This call returns \((27 \ 64 \ 8 \ 216)\).

Note that in this example, the first parameter to \texttt{mapcar} is a \texttt{LAMBDA} expression. When \texttt{EVAL} evaluates the \texttt{LAMBDA} expression, it constructs a function that has the same form as any predefined function except that it is nameless. In the example expression, this nameless function is immediately applied to each element of the parameter list and the results are returned in a list.

### 15.5.14 Functions That Build Code

The fact that programs and data have the same structure can be exploited in constructing programs. Recall that the Scheme interpreter uses a function named \texttt{EVAL}. The Scheme system applies \texttt{EVAL} to every expression typed, whether it is at the Scheme prompt in the interactive interpreter or is part of a program being interpreted. The \texttt{EVAL} function can also be called directly by Scheme programs. This provides the possibility of a Scheme program creating expressions and calling \texttt{EVAL} to evaluate them. This is not something that is unique to Scheme, but the simple forms of its expressions make it easy to create them during execution.

One of the simplest examples of this process involves numeric atoms. Recall that Scheme includes a function named \texttt{+}, which takes any number of numeric atoms as arguments and returns their sum. For example, \((+ \ 3 \ 7 \ 10 \ 2)\) returns \(22\).

Our problem is the following: Suppose that in a program we have a list of numeric atoms and need the sum. We cannot apply \texttt{+} directly on the list, because \texttt{+} can take only atomic parameters, not a list of numeric atoms. We could, of course, write a function that repeatedly adds the \texttt{CAR} of the list to the sum of its \texttt{CDR}, using recursion to go through the list. Such a function follows:

\[
\begin{align*}
\text{(DEFINE (adder a_list)} & \\
\text{(COND) } & \\
\text{((NULL? a_list) 0) } & \\
\text{(ELSE (+ (CAR a_list) (adder (CDR a_list))))})
\end{align*}
\]

Following is an example call to \texttt{adder}, along with the recursive calls and returns:

\[
\begin{align*}
\text{(adder '(3 4 5)) } & \\
\text{(+ 3 (adder (4 5))) } & \\
\text{(+ 3 (+ 4 (adder (5))))) } & \\
\text{(+ 3 (+ 4 (+ 5 (adder ())))) } & \\
\text{(+ 3 (+ 4 (+ 5 0)))) } & \\
\text{(+ 3 (+ 4 5)) } & \\
\text{(+ 3 9) } & \\
\text{(12) }
\end{align*}
\]

An alternative solution to the problem is to write a function that builds a call to \texttt{+} with the proper parameter forms. This can be done by using \texttt{CONS} to
build a new list that is identical to the parameter list except it has the atom `+` inserted at its beginning. This new list can then be submitted to `EVAL` for evaluation, as in the following:

```lisp
(DEFINE (adder a_list)
  (COND
    ((NULL? a_list) 0)
    (ELSE (EVAL (CONS '+ a_list)))
  ))
```

Note that the `+` function’s name is quoted to prevent `EVAL` from evaluating it in the evaluation of `CONS`. Following is an example call to this new version of `adder`, along with the call to `EVAL` and the return value:

```lisp
(adder '(3 4 5))
(EVAL (+ 3 4 5)
  (12)
```

In all earlier versions of Scheme, the `EVAL` function evaluated its expression in the outermost scope of the program. The later versions of Scheme, beginning with Scheme 4, requires a second parameter to `EVAL` that specifies the scope in which the expression is to be evaluated. For simplicity’s sake, we left the scope parameter out of our example, and we do not discuss scope names here.

## 15.6 Common Lisp

Common Lisp (Steele, 1990) was created in an effort to combine the features of several early 1980s dialects of Lisp, including Scheme, into a single language. Being something of a union of languages, it is quite large and complex, similar in these regards to C++ and C#. Its basis, however, is the original Lisp, so its syntax, primitive functions, and fundamental nature come from that language.

Following is the factorial function written in Common Lisp:

```lisp
(DEFUN factorial (x)
  (IF (<= n 1)
      1
      (* n factorial (- n 1)))
)
```

Only the first line of this function differs syntactically from the Scheme version of the same function.

The list of features of Common Lisp is long: a large number of data types and structures, including records, arrays, complex numbers, and character strings; powerful input and output operations; and a form of packages for modularizing collections of functions and data, and also for providing access control. Common Lisp includes several imperative constructs, as well as some mutable types.
Recognizing the occasional flexibility provided by dynamic scoping, as well as the simplicity of static scoping, Common Lisp allows both. The default scoping for variables is static, but by declaring a variable to be “special,” that variable becomes dynamically scoped.

Macros are often used in Common Lisp to extend the language. In fact, some of the predefined functions are actually macros. For example, DOLIST, which takes two parameters, a variable and a list, is a macro. For example, consider the following:

\[(\text{DOLIST} \ (x \ '(1 \ 2 \ 3)) \ \text{(print} \ x))\]

This produces the following:

```
1
2
3
NIL
```

NIL here is the return value of DOLIST.

Macros create their effect in two steps: First, the macro is expanded. Second, the expanded macro, which is Lisp code, is evaluated. Users can define their own macros with DEFMACRO.

The Common Lisp backquote operator (``) is similar to Scheme’s QUOTE, except some parts of the parameter can be unquoted by preceding them with commas. For example, consider the following two examples:

```
\'(a \ (* \ 3 \ 4) \ c)
```

This expression evaluates to \((a \ (* \ 3 \ 4) \ c)\). However, the following expression:

```
\'(a \ ,\ (* \ 3 \ 4) \ c)
```

evaluates to \((a \ 12 \ c)\).

Lisp implementations have a front end called the reader that transforms the text of Lisp programs into a code representation. Then, the macro calls in the code representation are expanded into code representations. The output of this step is then either interpreted or compiled into the machine language of the host computer, or perhaps into an intermediate code than can be interpreted. There is a special kind of macro, named reader macros or read macros, that are expanded during the reader phase of a Lisp language processor. A reader macro expands a specific character into a string of Lisp code. For example, the apostrophe in Lisp is a read macro that expands to a call to QUOTE. Users can define their own reader macros to create other shorthand constructs.

Common Lisp, as well as other Lisp-based languages, have a symbol data type. The reserved words are symbols that evaluate to themselves, as are \(\text{T}\) and \(\text{NIL}\). Technically, symbols are either bound or unbound. Parameter symbols are bound while the function is being evaluated. Also, symbols that are the
names of imperative-style variables and have been assigned values are bound. Other symbols are unbound. For example, consider the following expression:

\[(\text{LIST } '(\text{A B C}))\]

The symbols \text{A}, \text{B}, and \text{C} are unbound. Recall that Ruby also has a symbol data type.

In a sense, Scheme and Common Lisp are opposites. Scheme is far smaller and semantically simpler, in part because of its exclusive use of static scoping, but also because it was designed to be used for teaching programming, whereas Common Lisp was meant to be a commercial language. Common Lisp has succeeded in being a widely used language for AI applications, among other areas. Scheme, on the other hand, is more frequently used in college courses on functional programming. It is also more likely to be studied as a functional language because of its relatively small size. An important design goal of Common Lisp that caused it to be a large language was the desire to make it compatible with several of the dialects of Lisp from which it was derived.

The Common Lisp Object System (CLOS) (Paepeke, 1993) was developed in the late 1980s as an object-oriented version of Common Lisp. This language supports generic functions and multiple inheritance, among other constructs.

\section*{15.7 ML}

ML (Milner et al., 1997) is a static-scoped functional programming language, like Scheme. However, it differs from Lisp and its dialects, including Scheme, in a number of significant ways. One important difference is that ML is a strongly typed language, whereas Scheme is essentially typeless. ML has type declarations for function parameters and the return types of functions, although because of its type inferencing they are often not used. The type of every variable and expression can be statically determined. ML, like other functional programming languages, does not have variables in the sense of the imperative languages. It does have identifiers, which have the appearance of names of variables in imperative languages. However, these identifiers are best thought of as names for values. Once set, they cannot be changed. They are like the named constants of imperative languages like \text{final} declarations in Java. ML identifiers do not have fixed types—any identifier can be the name of a value of any type.

A table called the \textit{evaluation environment} stores the names of all implicitly and explicitly declared identifiers in a program, along with their types. This is like a run-time symbol table. When an identifier is declared, either implicitly or explicitly, it is placed in the evaluation environment.

Another important difference between Scheme and ML is that ML uses a syntax that is more closely related to that of an imperative language than that of Lisp. For example, arithmetic expressions are written in ML using infix notation.
Function declarations in ML appear in the general form

```ml
fun function_name(formal parameters) = expression;
```

When called, the value of the expression is returned by the function. Actually, the expression can be a list of expressions, separated by semicolons and surrounded by parentheses. The return value in this case is that of the last expression. Of course, unless they have side effects, the expressions before the last serve no purpose. Because we are not considering the parts of ML that have side effects, we only consider function definitions with a single expression.

Now we can discuss type inference. Consider the following ML function declaration:

```ml
fun circumf(r) = 3.14159 * r * r;
```

This specifies a function named `circumf` that takes a floating-point (real in ML) parameter and produces a floating-point result. The types are inferred from the type of the literal in the expression. Likewise, in the function

```ml
fun times10(x) = 10 * x;
```

the parameter and functional value are inferred to be of type int.

Consider the following ML function:

```ml
fun square(x) = x * x;
```

ML determines the type of both the parameter and the return value from the * operator in the function definition. Because this is an arithmetic operator, the type of the parameter and the function are assumed to be numeric. In ML, the default numeric type is int. So, it is inferred that the type of the parameter and the return value of `square` is int.

If `square` were called with a floating-point value, as in `square(2.75);` it would cause an error, because ML does not coerce real values to int type. If we wanted `square` to accept real parameters, it could be rewritten as

```ml
fun square(x) : real = x * x;
```

Because ML does not allow overloaded functions, this version could not coexist with the earlier int version. The last version defined would be the only one.

The fact that the functional value is typed real is sufficient to infer that the parameter is also real type. Each of the following definitions is also legal:

```ml
fun square(x : real) = x * x;
fun square(x) = (x : real) * x;
fun square(x) = x * (x : real);
```
Type inference is also used in the functional languages Miranda, Haskell, and F#.

The ML selection control flow construct is similar to that of the imperative languages. It has the following general form:

\[ \text{if expression then then_expression else else_expression} \]

The first expression must evaluate to a Boolean value.

The conditional expressions of Scheme can appear at the function definition level in ML. In Scheme, the `COND` function is used to determine the value of the given parameter, which in turn specifies the value returned by `COND`. In ML, the computation performed by a function can be defined for different forms of the given parameter. This feature is meant to mimic the form and meaning of conditional function definitions in mathematics. In ML, the particular expression that defines the return value of a function is chosen by pattern matching against the given parameter. For example, without using this pattern matching, a function to compute factorial could be written as follows:

\[
\begin{align*}
\text{fun fact}(n) : int = & \text{if } n \leq 1 \text{ then } 1 \\
& \text{else } n \times \text{fact}(n - 1);
\end{align*}
\]

Multiple definitions of a function can be written using parameter pattern matching. The different function definitions that depend on the form of the parameter are separated by an OR symbol (|). For example, using pattern matching, the factorial function could be written as follows:

\[
\begin{align*}
\text{fun fact}(0) = & 1 \\
| \text{fact}(1) = & 1 \\
| \text{fact}(n) : int = & n \times \text{fact}(n - 1);
\end{align*}
\]

If `fact` is called with the actual parameter 0, the first definition is used; if the actual parameter is 1, the second definition is used; if an `int` value that is neither 0 nor 1 is sent, the third definition is used.

As discussed in Chapter 6, ML supports lists and list operations. Recall that \( \text{hd}, \text{tl}, \text{and} \ :: \) are ML's versions of Scheme's \( \text{CAR}, \text{CDR}, \text{and} \ \text{CONS}. \)

Because of the availability of patterned function parameters, the \( \text{hd} \) and \( \text{tl} \) functions are much less frequently used in ML than \( \text{CAR} \) and \( \text{CDR} \) are used in Scheme. For example, in a formal parameter, the expression

\( (h :: t) \)

is actually two formal parameters, the head and tail of given list parameter, while the single corresponding actual parameter is a list. For example, the number of elements in a given list can be computed with the following function:

\[
\begin{align*}
\text{fun length}([]) = & 0 \\
| \text{length}(h :: t) = & 1 + \text{length}(t);
\end{align*}
\]
As another example of these concepts, consider the append function, which does what the Scheme append function does:

```haskell
fun append([], lis2) = lis2
| append(h :: t, lis2) = h :: append(t, lis2);
```

The first case in this function handles the situation of the function being called with an empty list as the first parameter. This case also terminates the recursion when the initial call has a nonempty first parameter. The second case of the function breaks the first parameter list into its head and tail (hd and tl). The head is CONSed onto the result of the recursive call, which uses the tail as its first parameter.

In ML, names are bound to values with value declaration statements of the form

```ml
val new_name = expression;
```

For example,

```ml
val distance = time * speed;
```

Do not get the idea that this statement is exactly like the assignment statements in the imperative languages, for it is not. The val statement binds a name to a value, but the name cannot be later rebound to a new value. Well, in a sense it can. Actually, if you do rebind a name with a second val statement, it causes a new entry in the evaluation environment that is not related to the previous version of the name. In fact, after the new binding, the old evaluation environment entry (for the previous binding) is no longer visible. Also, the type of the new binding need not be the same as that of the previous binding. val statements do not have side effects. They simply add a name to the current evaluation environment and bind it to a value.

The normal use of val is in a let expression. Consider the following example:

```ml
let val radius = 2.7
    val pi = 3.14159
in pi * radius * radius
end;
```

ML includes several higher-order functions that are commonly used in functional programming. Among these are a filtering function for lists, filter, which takes a predicate function as its parameter. The predicate function is often given as a lambda expression, which in ML is defined exactly like a function, except with the fn reserved word, instead of fun, and of course the lambda expression is nameless. filter returns a function that takes a list as a parameter. It tests each element of the list with the predicate. Each element on which the predicate returns true is added to a new list, which is the return value of the function. Consider the following use of filter:

9. let expressions in ML were introduced in Chapter 5.
filter(fn(x) => x < 100, [25, 1, 50, 711, 100, 150, 27, 161, 3]);

This application would return [25, 1, 50, 27, 3].

The map function takes a single parameter, which is a function. The resulting function takes a list as a parameter. It applies its function to each element of the list and returns a list of the results of those applications. Consider the following code:

```ml
fun cube x = x * x * x;
val cubeList = map cube;
val newList = cubeList [1, 3, 5];
```

After execution, the value of `newList` is [1, 27, 125]. This could be done more simply by defining the cube function as a lambda expression, as in the following:

```ml
val newList = map (fn x => x * x * x, [1, 3, 5]);
```

ML has a binary operator for composing two functions, o (a lowercase “oh”). For example, to build a function `h` that first applies function `f` and then applies function `g` to the returned value from `f`, we could use the following:

```ml
val h = g o f;
```

Strictly speaking, ML functions take a single parameter. When a function is defined with more than one parameter, ML considers the parameters to be a tuple, even though the parentheses that normally delimit a tuple value are optional. The commas that separate the parameters (tuple elements) are required.

The process of currying replaces a function with more than one parameter with a function with one parameter that returns a function that takes the other parameters of the initial function.

ML functions that take more than one parameter can be defined in curried form by leaving out the commas between the parameters (and the delimiting parentheses). For example, we could have the following:

```ml
fun add a b = a + b;
```

Although this appears to define a function with two parameters, it actually defines one with just one parameter. The `add` function takes an integer parameter (`a`) and returns a function that also takes an integer parameter (`b`). A call to this function also excludes the commas between the parameters, as in the following:

```ml
add 3 5;
```

This call to `add` returns 8, as expected.

---

10. This form of functions is named for Haskell Curry, a British mathematician who studied them.
Curried functions are interesting and useful because new functions can be constructed from them by partial evaluation. **Partial evaluation** means that the function is evaluated with actual parameters for one or more of the leftmost formal parameters. For example, we could define a new function as follows:

```plaintext
fun add5 x = add 5 x;
```

The `add5` function takes the actual parameter `5` and evaluates the `add` function with `5` as the value of its first formal parameter. It returns a function that adds `5` to its single parameter, as in the following:

```plaintext
val num = add5 10;
```

The value of `num` is now `15`. We could create any number of new functions from the curried function `add` to add any specific number to a given parameter.

Curried functions also can be written in Scheme, Haskell, and F#. Consider the following Scheme function:

```plaintext
(define (add x y) (+ x y))
```

A curried version of this would be as follows:

```plaintext
(define (add y) (lambda (x) (+ y x)))
```

This can be called as follows:

```plaintext
((add 3) 4)
```

ML has enumerated types, arrays, and tuples. ML also has exception handling and a module facility for implementing abstract data types.

ML has had a significant impact on the evolution of programming languages. For language researchers, it has become one of the most studied languages. Furthermore, it has spawned several subsequent languages, among them Haskell, Caml, OCaml, and F#.

### 15.8 Haskell

Haskell (Thompson, 1999) is similar to ML in that it uses a similar syntax, is static scoped, is strongly typed, and uses the same type inferencing method. There are three characteristics of Haskell that set it apart from ML: First, functions in Haskell can be overloaded (functions in ML cannot). Second, nonstrict semantics are used in Haskell, whereas in ML (and most other programming languages) strict semantics are used. Third, Haskell is a pure functional programming language, meaning it has no expressions or statements that have side effects, whereas ML allows some side effects (for example, ML has mutable...
arrays). Both nonstrict semantics and function overloading are further discussed later in this section.

The code in this section is written in version 1.4 of Haskell.

Consider the following definition of the factorial function, which uses pattern matching on its parameters:

```haskell
fact 0 = 1
fact 1 = 1
fact n = n * fact (n - 1)
```

Note the differences in syntax between this definition and its ML version in Section 15.7. First, there is no reserved word to introduce the function definition (`fun` in ML). Second, alternative definitions of functions (with different formal parameters) all have the same appearance.

Using pattern matching, we can define a function for computing the $n$th Fibonacci number with the following:

```haskell
fib 0 = 1
fib 1 = 1
fib (n + 2) = fib (n + 1) + fib n
```

Guards can be added to lines of a function definition to specify the circumstances under which the definition can be applied. For example,

```haskell
fact n
    | n == 0 = 1
    | n == 1 = 1
    | n > 1 = n * fact(n - 1)
```

This definition of factorial is more precise than the previous one, for it restricts the range of actual parameter values to those for which it works. This form of a function definition is called a *conditional expression*, after the mathematical expressions on which it is based.

An *otherwise* can appear as the last condition in a conditional expression, with the obvious semantics. For example,

```haskell
sub n
    | n < 10 = 0
    | n > 100 = 2
    | otherwise = 1
```

Notice the similarity between the guards here and the guarded commands discussed in Chapter 8.

Consider the following function definition, whose purpose is the same as the corresponding ML function in Section 15.7:

```haskell
square x = x * x
```
In this case, however, because of Haskell’s support for polymorphism, this function can take a parameter of any numeric type.

As with ML, lists are written in brackets in Haskell, as in

```haskell
colors = ["blue", "green", "red", "yellow"]
```

Haskell includes a collection of list operators. For example, lists can be catenated with \+\+, : serves as an infix version of \texttt{CONS}, and \ldots\ is used to specify an arithmetic series in a list. For example,

\[
\begin{align*}
5: &[2, 7, 9] & \text{ results in } & [5, 2, 7, 9] \\
[1, 3..11] & \text{ results in } & [1, 3, 5, 7, 9, 11] \\
[1, 3, 5] & \text{} + [2, 4, 6] & \text{ results in } & [1, 3, 5, 2, 4, 6]
\end{align*}
\]

Notice that the : operator is just like ML’s : operator.\footnote{It is interesting that ML uses : for attaching a type name to a name and :: for \texttt{CONS}, while Haskell uses these two operators in exactly opposite ways.} Using : and pattern matching, we can define a simple function to compute the product of a given list of numbers:

```haskell
product [] = 1
product (a:x) = a * product x
```

Using \texttt{product}, we can write a factorial function in the simpler form

\[
\text{fact } n = \text{product } [1..n]
\]

Haskell includes a \texttt{let} construct that is similar to ML’s \texttt{let} and \texttt{val}. For example, we could write

```haskell
quadratic_root a b c =
  let minus_b_over_2a = - b / (2.0 * a)
      root_part_over_2a =
        sqrt(b ^ 2 - 4.0 * a * c) / (2.0 * a)
    in
      minus_b_over_2a - root_part_over_2a,
      minus_b_over_2a + root_part_over_2a
```

Haskell’s list comprehensions were introduced in Chapter 6. For example, consider the following:

\[
[n * n * n | n <- [1..50]]
\]

This defines a list of the cubes of the numbers from 1 to 50. It is read as “a list of all $n^n*n$ such that $n$ is taken from the range of 1 to 50.” In this case, the qualifier is in the form of a generator. It generates the numbers from 1 to 50. In other
cases, the qualifiers are in the form of Boolean expressions and they are called **tests**. This notation can be used to describe algorithms for doing many things, such as finding permutations of lists and sorting lists. For example, consider the following function, which when given a number \( n \) returns a list of all its factors:

\[
\text{factors } n = \{ i \mid i \leftarrow [1..n \ `\text{div}` 2], n \ `\text{mod}` i == 0 \}
\]

The list comprehension in `factors` creates a list of numbers, each temporarily bound to the name `i`, ranging from 1 to \( n/2 \), such that \( n \ `\text{mod}` i \) is zero. This is indeed a very exacting and short definition of the factors of a given number. The backticks (backward apostrophes) surrounding `div` and `mod` are used to specify the infix use of these functions. When they are called in functional notation, as in `div n 2`, the backticks are not used.

Next, consider the concision of Haskell illustrated in the following implementation of the quicksort algorithm:

\[
\text{sort } [] = []
\]
\[
\text{sort } (h:t) = \text{sort } [b \mid b \leftarrow t, b \leftarrow h] ++ \{h\} ++ \text{sort } [b \mid b \leftarrow t, b > h]
\]

In this program, the set of list elements that are smaller or equal to the list head are sorted and catenated with the head element, then the set of elements that are greater than the list head are sorted and catenated onto the previous result. This definition of quicksort is significantly shorter and simpler than the same algorithm coded in an imperative language.

A programming language is **strict** if it requires all actual parameters to be fully evaluated, which ensures that the value of a function does not depend on the order in which the parameters are evaluated. A language is **nonstrict** if it does not have the strict requirement. Nonstrict languages can have several distinct advantages over strict languages. First, nonstrict languages are generally more efficient, because some evaluation is avoided. Second, some interesting capabilities are possible with nonstrict languages that are not possible with strict languages. Among these are infinite lists. Nonstrict languages can use an evaluation form called **lazy evaluation**, which means that expressions are evaluated only if and when their values are needed.

Recall that in Scheme the parameters to a function are fully evaluated before the function is called, so it has strict semantics. Lazy evaluation means that an actual parameter is evaluated only when its value is necessary to evaluate the function. So, if a function has two parameters, but on a particular execution of the function the first parameter is not used, the actual parameter passed for that execution will not be evaluated. Furthermore, if only a part of an actual parameter must be evaluated for an execution of the function, the rest is left

---

12. Notice how this is related to short-circuit evaluation of Boolean expressions, which is done in some imperative languages.
unevaluated. Finally, actual parameters are evaluated only once, if at all, even if the same actual parameter appears more than once in a function call.

As stated previously, lazy evaluation allows one to define infinite data structures. For example, consider the following:

```plaintext
positives = [0..]
evens = [2, 4..]
squares = [n * n | n <- [0..]]
```

Of course, no computer can actually represent all of the numbers of these lists, but that does not prevent their use if lazy evaluation is used. For example, if we wanted to know if a particular number was a perfect square, we could check the `squares` list with a membership function. Suppose we had a predicate function named `member` that determined whether a given atom is contained a given list. Then we could use it as in

```plaintext
member 16 squares
```

which would return `True`. The `squares` definition would be evaluated until the `16` was found. The `member` function would need to be carefully written. Specifically, suppose it were defined as follows:

```plaintext
member b [] = False
member b (a:x)= (a == b) || member b x
```

The second line of this definition breaks the first parameter into its head and tail. Its return value is true if either the head matches the element for which it is searching (`b`) or if the recursive call with the tail of the list returns `True`.

This definition of `member` would work correctly with `squares` only if the given number were a perfect square. If not, `squares` would keep generating squares forever, or until some memory limitation was reached, looking for the given number in the list. The following function performs the membership test of an ordered list, abandoning the search and returning `False` if a number greater than the searched-for number is found.¹³

```plaintext
member2 n (m:x)
| m < n     = member2 n x
| m == n    = True
| otherwise = False
```

Lazy evaluation sometimes provides a modularization tool. Suppose that in a program there is a call to function `f` and the parameter to `f` is the return value of a function `g`.¹⁴ So, we have `f (g (x))`. Further suppose that `g` produces a large amount of data, a little at a time, and that `f` must then process this data,

---

¹³. This assumes that the list is in ascending order.

¹⁴. This example appears in Hughes (1989).
a little at a time. In a conventional imperative language, \( g \) would run on the whole input producing a file of its output. Then \( f \) would run using the file as its input. This approach requires the time to both write and read the file, as well as the storage for the file. With lazy evaluation, the executions of \( f \) and \( g \) implicitly would be tightly synchronized. Function \( g \) will execute only long enough to produce enough data for \( f \) to begin its processing. When \( f \) is ready for more data, \( g \) will be restarted to produce more, while \( f \) waits. If \( f \) terminates without getting all of \( g \)'s output, \( g \) is aborted, thereby avoiding useless computation. Also, \( g \) need not be a terminating function, perhaps because it produces an infinite amount of output. \( g \) will be forced to terminate when \( f \) terminates. So, under lazy evaluation, \( g \) runs as little as possible. This evaluation process supports the modularization of programs into generator units and selector units, where the generator produces a large number of possible results and the selector chooses the appropriate subset.

Lazy evaluation is not without its costs. It would certainly be surprising if such expressive power and flexibility were free. In this case, the cost is in a far more complicated semantics, which results in much slower speed of execution.

### 15.9 F#

F# is a .NET functional programming language whose core is based on OCaml, which is a descendant of ML and Haskell. Although it is fundamentally a functional language, it includes imperative features and supports object-oriented programming. One of the most important characteristics of F# is that it has a full-featured IDE, an extensive library of utilities that supports imperative, object-oriented, and functional programming, and has interoperability with a collection of nonfunctional languages (all of the .NET languages).

F# is a first-class .NET language. This means that F# programs can interact in every way with other .NET languages. For example, F# classes can be used and subclassed by programs in other languages, and vice versa. Furthermore, F# programs have access to all of the .NET Framework APIs. The F# implementation is available free from Microsoft ([http://research.microsoft.com/fsharp/fsharp.aspx](http://research.microsoft.com/fsharp/fsharp.aspx)). It is also supported by Visual Studio.

F# includes a variety of data types. Among these are tuples, like those of Python and the functional languages ML and Haskell, lists, discriminated unions, an expansion of ML's unions, and records, like those of ML, which are like tuples except the components are named. F# has both mutable and immutable arrays.

Recall from Chapter 6, that F#’s lists are similar to those of ML, except that the elements are separated by semicolons and \( \text{hd} \) and \( \text{tl} \) must be called as methods of \text{List}.

F# supports sequence values, which are types from the .NET namespace \text{System.Collections.Generic(IEnumerable)}. In F#, sequences are abbreviated as \text{seq<type>}, where \text{<type>} indicates the type of the generic. For example, the type \text{seq<int>} is a sequence of integer values. Sequence values
can be created with generators and they can be iterated. The simplest sequences are generated with range expressions, as in the following example:

```fsharp
let x = seq {1..4};;
```

In the examples of F#, we assume that the interactive interpreter is used, which requires the two semicolons at the end of each statement. The expression above generates `seq[1; 2; 3; 4]`. (List and sequence elements are separated by semicolons.) The generation of a sequence is lazy; for example, the following defines `y` to be a very long sequence, but only the needed elements are generated. For display, only the first four are generated.

```fsharp
let y = seq {0..100000000};;
y;;
val it: seq<int> = seq[0; 1; 2; 3; . . .]
```

The first line above defines `y`; the second line requests that the value of `y` be displayed; the third is the output of the F# interactive interpreter.

The default step size for integer sequence definitions is `1`, but it can be set by including it in the middle of the range specification, as in the following example:

```fsharp
seq {1..2..7};;
```

This generates `seq [1; 3; 5; 7]`.

The values of a sequence can be iterated with a `for-in` construct, as in the following example:

```fsharp
let seq1 = seq {0..3..11};;
for value in seq1 do printfn "value = %d" value;;
```

This produces the following:

```
value = 0
value = 3
value = 6
value = 9
```

Iterators can also be used to create sequences, as in the following example:

```fsharp
let cubes = seq {
    for i in 1..5 -> (i, i * i * i)};;
```

This generates the following list of tuples:

```
seq [(1, 1); (2, 8); (3, 27); (4, 64); (5, 125)]
```

This use of iterators to generate collections is a form of list comprehension.
Sequencing can also be used to generate lists and arrays, although in these cases the generation is not lazy. In fact, the primary difference between lists and sequences in F# is that sequences are lazy, and thus can be infinite, whereas lists are not lazy. Lists are in their entirety stored in memory. That is not the case with sequences.

The functions of F# are similar to those of ML and Haskell. If named, they are defined with let statements. If unnamed, which means technically they are lambda expressions, they are defined with the fun reserved word. The following lambda expression illustrates their syntax:

\[
\text{fun } \ a \ b \to a / b
\]

Note that there is no difference between a name defined with let and a function without parameters defined with let.

Indentation is used to show the extent of a function definition. For example, consider the following function definition:

\[
\text{let } f = \\
\quad \text{let } \pi = 3.14159 \\
\quad \text{let } \text{twoPi} = 2.0 \times \pi \\
\quad \text{twoPi};;
\]

Note that F#, like ML, does not coerce numeric values, so if this function used 2 in the second-last line, rather than 2.0, an error would be reported.

If a function is recursive, the reserved word rec must precede its name in its definition. Following is an F# version of factorial:

\[
\text{let rec } \text{factorial} \ x = \\
\quad \text{if } x \leq 1 \ \text{then} \ 1 \\
\quad \text{else } n \times \text{factorial}(n - 1)
\]

Names defined in functions can be outscoped, which means they can be redefined, which ends their former scope. For example, we could have the following:

\[
\text{let } x^4 \ x = \\
\quad \text{let } \ x = \ x \times \ x \\
\quad \text{let } \ x = \ x \times \ x \\
\quad \ x;;
\]

In this function, the first let in the body of the \(x^4\) function creates a new version of \(x\), defining it to have the value of the square of the parameter \(x\). This terminates the scope of the parameter. So, the second let in the function body uses the new \(x\) in its right side and creates yet another version of \(x\), thereby terminating the scope of the \(x\) created in the previous let.

There are two important functional operators in F#, pipeline (\(\mid\)) and function composition (\(\circ\)). The pipeline operator is a binary operator that
sends the value of its left operand, which is an expression, to the last parameter of the function call, which is the right operand. It is used to chain together function calls while flowing the data being processed to each call. Consider the following example code, which uses the high-order functions \texttt{filter} and \texttt{map}:

```fsharp
let myNums = [1; 2; 3; 4; 5]
let evensTimesFive = myNums
    |> List.filter (fun n -> n % 2 = 0)
    |> List.map (fun n -> 5 * n)
```

The \texttt{evensTimesFive} function begins with the list \texttt{myNums}, filters out the numbers that are not even with \texttt{filter}, and uses \texttt{map} to map a lambda expression that multiplies the numbers in a given list by five. The return value of \texttt{evensTimesFive} is \([10; 20]\).

The function composition operator builds a function that applies its left operand to a given parameter, which is a function, and then passes the result returned from that function to its right operand, which is also a function. So, the F# expression \((f >> g) x\) is equivalent to the mathematical expression \(g(f(x))\).

Like ML, F# supports curried functions and partial evaluation. The ML example in Section 15.7 could be written in F# as follows:

```fsharp
let add a b = a + b;;
let add5 = add 5;;
```

Note that, unlike ML, the syntax of the formal parameter list in F# is the same for all functions, so all functions with more than one parameter can be curried.

F# is interesting for several reasons: First, it builds on the past functional languages as a functional language. Second, it supports virtually all programming methodologies in widespread use today. Third, it is the first functional language that is designed for interoperability with other widely used languages. Fourth, it starts out with an elaborate and well-developed IDE and library of utility software with .NET and its framework.

### 15.10 Support for Functional Programming in Primarily Imperative Languages

Imperative programming languages have typically provided only limited support for functional programming. That limited support has resulted in little use of those languages for functional programming. The most important restriction, related to functional programming, of imperative languages of the past was the lack of support for higher-order functions.

One indication of the increasing interest and use of functional programming is the partial support for it that has begun to appear over the last decade in programming languages that are primarily imperative. For example,
anonymous functions, which are like lambda expressions, are now part of JavaScript, Python, Ruby, Java, and C#.

In JavaScript, named functions are defined with the following syntax:

```javascript
function name (formal-parameters) {
    body
}
```

An anonymous function is defined in JavaScript with the same syntax, except that the name of the function is omitted.

In C#, a lambda expression is an instance of a delegate. They can be anonymous or named. An anonymous lambda expression is simpler than an anonymous method because methods must define their parameter types and return type, but lambda expressions use the C# inferencing process to avoid those necessities. The syntax of an unnamed lambda expression in C# is as follows:

```csharp
parameter(s) => expression
```

If there is more than one parameter, they must be enclosed in parentheses. If there are no parameters, empty parentheses must appear in the place of the parameters. If the system cannot infer the types of the parameters, they can be preceded by type names. The type of the return value is never specified; it is always inferred by the context of the lambda expression. The expression is either a single expression or a compound statement enclosed in braces. Such a compound statement must include a `return` statement.

One common use of anonymous lambda expressions is as actual parameters to methods that are specified to take delegates as parameters. For example, C# has a collection of methods for arrays that perform searching operations. For example, the `FindAll` method finds all of the elements of an array that satisfy a given instance of a delegate that takes a parameter of the type of the array's elements and returns a Boolean value. For example, we could have the following:

```csharp
int[] numbers = {-3, 0, 4, 5, 1, 7, -3, -6, -9, 0, 3};
int[] positives = Array.FindAll(numbers, n => n > 0);
// Now, positives is {4, 5, 1, 7, 3}
```

Lambda expressions in C# can also be named. The language has generic delegates that make defining such lambda expressions simple, although they do not cover all possibilities. One commonly used generic delegate is `Func`, which can take up to sixteen generic parameters for the parameters of the lambda expression, plus one more for the return type. For example, consider the following example of a named lambda expression and an invocation of it:

```csharp
Func<int, int, int> eval1 = (a, b) => 3 * a + (b / 2);
int result = eval1(6, 22);
```

C# lambda expressions can access variables defined outside their definitions. When they do, the lifetimes of the accessed external variables, which are called
captured variables, are extended so that they still exist when the lambda expression is used. A lambda expression that captures outside variables is a closure.

Lambda expressions were added to Java 8. The general syntax of these expressions is like that of C#, except that \(-\rightarrow\) is used instead of \(\Rightarrow\). The syntax of parameters, the inferencing of parameter types and the return type, and the expression or block with a return are the same as with C#. Prior to Java 8, there was no convenient way to pass a block of code to a method or have the method return a block of code.\(^{15}\)

Python’s lambda expressions define simple one-statement anonymous functions. The syntax of a lambda expression in Python is exemplified by the following:

\[
\text{lambda } a, b : 2 * a - b
\]

Note that the formal parameters are separated from function body by a colon.

Python includes the higher-order functions \texttt{filter} and \texttt{map}. Both often use lambda expressions as their first parameter. The second parameter of these is a sequence type, and both return the same sequence type as their second parameter. In Python, strings, lists, and tuples are considered sequences. Consider the following example of using the map function in Python:

\[
\text{map(}\text{lambda } x: x ** 3, \{2, 4, 6, 8\})
\]

This call returns \([8, 64, 216, 512]\).

Python also supports partial function applications. Consider the following example:

\[
\text{from operator import add} \\
\text{add5 = partial (add, 5)}
\]

The \texttt{from} declaration here imports the functional version of the addition operator, which is named \texttt{add}, from the \texttt{operator} module.

After defining \texttt{add5}, it can be used with one parameter, as in the following:

\[
\text{add5(15)}
\]

This call returns \(20\).

As described in Chapter 6, Python includes lists and list comprehensions.

Ruby’s blocks are effectively subprograms that are sent to methods, which makes the method a higher-order subprogram. A Ruby block can be converted to a subprogram object with \texttt{lambda}. For example, consider the following:

\[
\text{times = lambda \{ |a, b| a * b\}}
\]

\(^{15}\). One inconvenient way was to define a class with a method that contained the code, instantiate the class, and pass a reference to it.
Following is an example of using `times`:

\[ x = \text{times}.(3, 4) \]

This sets \( x \) to 12. The `times` object can be curried with the following:

\[ \text{times}5 = \text{times}.\text{curry}.(5) \]

This function can be used as in the following:

\[ x5 = \text{times}5.(3) \]

This sets \( x5 \) to 15.

### 15.11 A Comparison of Functional and Imperative Languages

This section discusses some of the differences between imperative and functional languages.

Functional languages can have a very simple syntactic structure. The list structure of Lisp, which is used for both code and data, clearly illustrates this. The syntax of the imperative languages is much more complex. This makes them more difficult to learn and to use.

The semantics of functional languages is also simpler than that of the imperative languages. For example, in the denotational semantics description of an imperative loop construct given in Section 3.5.2, the loop is converted from an iterative construct to a recursive construct. This conversion is unnecessary in a pure functional language, in which there is no iteration. Furthermore, we assumed there were no expression side effects in all of the denotational semantic descriptions of imperative constructs in Chapter 3. This restriction is unrealistic for imperative languages, because all of the C-based languages include expression side effects. This restriction is not needed for the denotational descriptions of pure functional languages.

Some in the functional programming community have claimed that the use of functional programming results in an order-of-magnitude increase in productivity, largely due to functional programs being claimed to be only 10 percent as large as their imperative counterparts. While such numbers have been actually shown for certain problem areas, for other problem areas, functional programs are more like 25 percent as large as imperative solutions to the same problems (Wadler, 1998). These factors allow proponents of functional programming to claim productivity advantages over imperative programming of 4 to 10 times. However, program size alone is not necessarily a good measure of productivity. Certainly not all lines of source code have equal complexity, nor do they take the same amount of time to produce. In fact, because of the necessity of dealing with variables, imperative programs have many trivially simple lines for initializing and making small changes to variables.
Execution efficiency is another basis for comparison. When functional programs are interpreted, they are of course much slower than their compiled imperative counterparts. However, there are now compilers for most functional languages, so that execution speed disparities between functional languages and compiled imperative languages are no longer so great. One might be tempted to say that because functional programs are significantly smaller than equivalent imperative programs, they should execute much faster than the imperative programs. However, this often is not the case, because of a collection of language characteristics of the functional languages, such as lazy evaluation, that have a negative impact on execution efficiency. Considering the relative efficiency of functional and imperative programs, it is reasonable to estimate that an average functional program will execute in about twice the time of its imperative counterpart (Wadler, 1998). This may sound like a significant difference, one that would often lead one to dismiss the functional languages for a given application. However, this factor-of-two difference is important only in situations where execution speed is of the utmost importance. There are many situations where a factor of two in execution speed is not considered important. For example, consider that many programs written in imperative languages, such as the Web software written in JavaScript and PHP, are interpreted and therefore are much slower than equivalent compiled versions. For these applications, execution speed is not the first priority.

Another source of the difference in execution efficiency between functional and imperative programs is the fact that imperative languages were designed to run efficiently on von Neumann architecture computers, while the design of functional languages is based on mathematical functions. This gives the imperative languages a large advantage.

Functional languages have a potential advantage in readability. In many imperative programs, the details of dealing with variables obscure the logic of the program. Consider a function that computes the sum of the cubes of the first \( n \) positive integers. In C, such a function would likely appear similar to the following:

```c
int sum_cubes(int n){
    int sum = 0;
    for(int index = 1; index <= n; index++)
        sum += index * index * index;
    return sum;
}
```

In Haskell, the function could be:

```haskell
sumCubes n = sum (map (^3) [1..n])
```

This version simply specifies three steps:

1. Build the list of numbers ([1..n]).
2. Create a new list by mapping a function that computes the cube of a number onto each number in the list.
3. Sum the new list.
Because of the lack of details of variables and iteration control, this version is more readable than the C version.¹⁶

Concurrent execution in the imperative languages is difficult to design and difficult to use, as we saw in Chapter 13. In an imperative language, the programmer must make a static division of the program into its concurrent parts, which are then written as tasks, whose execution often must be synchronized. This can be a complicated process. Programs in functional languages are naturally divided into functions. In a pure functional language, these functions are independent in the sense that they do not create side effects and their operations do not depend on any nonlocal or global variables. Therefore, it is much easier to determine which of them can be concurrently executed. The actual parameter expressions in calls often can be evaluated concurrently. Simply by specifying that it can be done, a function can be implicitly evaluated in a separate thread, as in Multilisp. And, of course, access to shared immutable data does not require synchronization.

One simple factor that strongly affects the complexity of imperative, or procedural programming, is the necessary attention of the programmer to the state of the program at each step of its development. In a large program, the state of the program is a large number of values (for the large number of program variables). In pure functional programming, there is no state; hence, no need to devote attention to keeping it in mind.

It is not a simple matter to determine precisely why functional languages have not attained greater popularity. The inefficiency of the early implementations was clearly a factor then, and it is likely that at least some contemporary imperative programmers still believe that programs written in functional languages are slow. In addition, the vast majority of programmers learn programming using imperative languages, which makes functional programs appear to them to be strange and difficult to understand. For many who are comfortable with imperative programming, the switch to functional programming is an unattractive and potentially difficult move. On the other hand, those who begin with a functional language never notice anything strange about functional programs.

**Summary**

Mathematical functions are named or unnamed mappings that use only conditional expressions and recursion to control their evaluations. Complex functions can be defined using higher-order functions or functional forms, in which functions are used as parameters, returned values, or both.

Functional programming languages are modeled on mathematical functions. In their pure form, they do not use variables or assignment statements to

¹⁶ Of course, the C version could have been written in a more functional style, but most C programmers probably would not write it that way.
produce results; rather, they use function applications, conditional expressions, and recursion for execution control and functional forms to construct complex functions. Lisp began as a purely functional language but soon acquired a number of imperative-language features added in order to increase its efficiency and ease of use.

The first version of Lisp grew out of the need for a list-processing language for AI applications. Lisp is still the most widely used language for that application area.

The first implementation of Lisp was serendipitous: The original version of `EVAL` was developed solely to demonstrate that a universal Lisp function could be written.

Because Lisp data and Lisp programs have the same form, it is possible to have a program build another program. The availability of `EVAL` allows dynamically constructed programs to be executed immediately.

Scheme is a relatively simple dialect of Lisp that uses static scoping exclusively. Like Lisp, Scheme’s primary primitives include functions for constructing and dismantling lists, functions for conditional expressions, and simple predicates for numbers, symbols, and lists.

Common Lisp is a Lisp-based language that was designed to include most of the features of the Lisp dialects of the early 1980s. It allows both static- and dynamic-scoped variables and includes many imperative features. Common Lisp uses macros to define some of its functions. Users are allowed to define their own macros. The language includes reader macros, which are also user definable. Reader macros define single-symbol macros.

ML is a static-scoped and strongly typed functional programming language that uses a syntax that is more closely related to that of an imperative language than to Lisp. It includes a type-inferencing system, exception handling, a variety of data structures, and abstract data types.

ML does not do any type coercions and does not allow function overloading. Multiple definitions of functions can be defined using pattern matching of the actual parameter form. Currying is the process of replacing a function that takes multiple parameters with one that takes a single parameter and returns a function that takes the other parameters. ML, as well as several other functional languages, supports currying.

Haskell is similar to ML, except that all expressions in Haskell are evaluated using a lazy method, which allows programs to deal with infinite lists. Haskell also supports list comprehensions, which provide a convenient and familiar syntax for describing sets. Unlike ML and Scheme, Haskell is a pure functional language.

F# is a .NET programming language that supports functional and imperative programming, including object-oriented programming. Its functional programming core is based on OCaml, a descendent of ML and Haskell. F# is supported by an elaborate and widely used IDE. It also interoperates with other .NET languages and has access to the .NET class library.
BIBLIOGRAPHIC NOTES

The first published version of Lisp can be found in McCarthy (1960). A widely used version from the mid-1960s until the late 1970s is described in McCarthy et al. (1965) and Weissman (1967). Common Lisp is described in Steele (1990). The Scheme language is described in Dybvig (2009). ML is defined in Milner et al. (1997). Ullman (1998) is an excellent introductory textbook for ML. Programming in Haskell is introduced in Thompson (1999). F# is described in Syme et al. (2010).

The Scheme programs in this chapter were developed using DrRacket’s legacy language R5RS.

A rigorous discussion of functional programming in general can be found in Henderson (1980). The process of implementing functional languages through graph reduction is discussed in detail in Peyton Jones (1987).

REVIEW QUESTIONS

1. Define functional form, simple list, bound variable, and referential transparency.
2. What does a lambda expression specify?
3. What data types were parts of the original Lisp?
4. In what common data structure are Lisp lists normally stored?
5. Explain why QUOTE is needed for a parameter that is a data list.
6. What is a simple list?
7. What does the abbreviation REPL stand for?
8. What are the three parameters to IF?
9. What are the differences between =, EQ?, EQV?, and EQUAL?
10. What are the differences between the evaluation method used for the Scheme special form DEFINE and that used for its primitive functions?
11. What are the two forms of DEFINE?
12. Describe the syntax and semantics of COND.
13. Why are CAR and CDR so named?
14. If CONS is called with two atoms, say ’A and ’B, what is the returned?
15. Describe the syntax and semantics of LET in Scheme.
16. What are the differences between CONS, LIST, and APPEND?
17. Describe the syntax and semantics of mapcar in Scheme.
18. What is tail recursion? Why is it important to define functions that use recursion to specify repetition to be tail recursive?
19. Why were imperative features added to most dialects of Lisp?
20. In what ways are Common Lisp and Scheme opposites?
21. What scoping rule is used in Scheme? In Common Lisp? In ML? In Haskell? In F#?
22. What happens during the reader phase of a Common Lisp language processor?
23. What are two ways that ML is fundamentally different from Scheme?
24. What is stored in an ML evaluation environment?
25. What is the difference between an ML val statement and an assignment statement in C?
26. What is type inferencing, as used in ML?
27. What is the use of the fn reserved word in ML?
28. Can ML functions that deal with scalar numerics be generic?
29. What is a curried function?
30. What does partial evaluation mean?
31. Describe the actions of the ML filter function.
32. What operator does ML use for Scheme’s CAR?
33. What operator does ML use for functional composition?
34. What are the three characteristics of Haskell that make it different from ML?
35. What does lazy evaluation mean?
36. What is a strict programming language?
37. What programming paradigms are supported by F#?
38. With what other programming languages can F# interoperate?
39. What does F#’s let do?
40. How is the scope of a F# let construct terminated?
41. What is the underlying difference between a sequence and a list in F#?
42. What is the difference between the let of ML and that of F#, in terms of extent?
43. What is the syntax of a lambda expression in F#?
44. Does F# coerce numeric values in expressions? Argue in support of the design choice.
45. What support does Python provide for functional programming?
46. What function in Ruby is used to create a curried function?
47. Is the use of functional programming expanding or shrinking?
48. What is one characteristic of functional programming languages that makes their semantics simpler than that of imperative languages?
49. What is the flaw in using lines of code to compare the productivity of functional languages and that of imperative languages?
50. Why can concurrency be easier with functional languages than imperative languages?
1. Read Robert Wilensky’s *Common LISPcraft* (Wilensky, 1986) and compare the features of Scheme discussed in this chapter with the corresponding features of LISP.

2. Find definitions of the Scheme functions `EVAL` and `APPLY`, and explain their actions.

3. One of the most modern and complete programming environments is the INTERLISP system for Lisp, as described in “The INTERLISP Programming Environment,” by Teitelmen and Masinter (*IEEE Computer*, Vol. 14, No. 4, April 1981). Read this article carefully and compare the difficulty of writing Lisp programs on your system with that of using INTERLISP (assuming that you do not normally use INTERLISP).

4. Refer to a book on Haskell programming and discuss the difference between ML and Haskell on function overloading.

5. Find at least one example of a typed functional programming language being used to build a commercial system in each of the following areas: telecommunication and embedded software.

6. A functional language could use some data structure other than the list. For example, it could use strings of single-character symbols. What primitives would such a language have in place of the `CAR`, `CDR`, and `CONS` primitives of Scheme?

7. Make a list of the features of F# that are not in ML.

8. If Scheme were a pure functional language, could it include `DISPLAY`? Why or why not?

9. What does the following ML function do?

   ```ml
   fun length ([ ]) = 0
   | length (h :: t) = 1 + length (t);
   ```

10. What does the following Scheme function do?

    ```scheme
    (define (x lis)
      (cond
        ((null? lis) 0)
        ((not (list? (car lis)))
          (cond
            ((eq? (car lis) #f) (x (cdr lis)))
            (else (+ 1 (x (cdr lis))))))
        (else (+ (x (car lis)) (x (cdr lis))))))
    ```
1. Write a Scheme function that computes the area of a circle, given its radius.

2. Write a Scheme function that computes the real roots of a given quadratic equation. If the roots are complex, the function must display a message indicating that. This function must use an IF function. The three parameters to the function are the three coefficients of the quadratic equation.

3. Repeat Programming Exercise 2 using a COND function, rather than an IF function.

4. Write a Scheme function that computes the factorial of a number.

5. Write an ML function that returns the cube of a given number.

6. Write a Scheme function that takes a simple list of numbers as a parameter and returns a list with the largest and smallest numbers in the input list.

7. Write a Scheme function that takes a list and an atom as parameters and returns a list identical to its parameter list except with all top-level instances of the given atom deleted.

8. Write a Scheme function that takes a list as a parameter and returns a list identical to the parameter except the last element has been deleted.

9. Repeat Programming Exercise 7, except that the atom can be either an atom or a list.

10. Write a Scheme function that takes two atoms and a list as parameters and returns a list identical to the parameter list except all occurrences of the first given atom in the list are replaced with the second given atom, no matter how deeply the first atom is nested.

11. Write a Scheme function that returns the reverse of its simple list parameter.

12. Write a Scheme predicate function that tests for the structural equality of two given lists. Two lists are structurally equal if they have the same list structure, although their atoms may be different.

13. Write a Scheme function that returns the union of two simple list parameters that represent sets.

14. Write a Scheme function with two parameters, an atom and a list, that returns a list identical to the parameter list except with all occurrences, no matter how deep, of the given atom deleted. The returned list cannot contain anything in place of the deleted atoms.

15. Write a Scheme function that takes a list as a parameter and returns a list identical to the parameter list except with the second top-level element removed. If the given list does not have two elements, the function should return ( ).
16. Write a Scheme function that takes a simple list of numbers as its parameter and returns a list identical to the parameter list except with the numbers in ascending order.

17. Write a Scheme function that takes a simple list of numbers as its parameter and returns the largest and smallest numbers in the list.

18. Write a Scheme function that takes a simple list as its parameter and returns a list of all permutations of the given list.

19. Write the quicksort algorithm in Scheme.

20. Rewrite the following Scheme function as a tail-recursive function:

   (DEFINE (doit n)
            (IF (= n 0)
                0
                (+ n (doit (- n 1)))
            ))

21. Write any of the first 19 Programming Exercises in F#.

22. Write any of the first 19 Programming Exercises in ML.
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Logic Programming Languages

16.1 Introduction
16.2 A Brief Introduction to Predicate Calculus
16.3 Predicate Calculus and Proving Theorems
16.4 An Overview of Logic Programming
16.5 The Origins of Prolog
16.6 The Basic Elements of Prolog
16.7 Deficiencies of Prolog
16.8 Applications of Logic Programming
Chapter 16  Logic Programming Languages

The objectives of this chapter are to introduce the concepts of logic programming and logic programming languages, including a brief description of a subset of Prolog. We begin with an introduction to predicate calculus, which is the basis for logic programming languages. This is followed by a discussion of how predicate calculus can be used for automatic theorem-proving systems. Then, we present a general overview of logic programming. Next, a lengthy section introduces the basics of the Prolog programming language, including arithmetic, list processing, and a trace tool that can be used to help debug programs and also to illustrate how the Prolog system works. The final two sections describe some of the problems of Prolog as a logic language and some of the application areas in which Prolog has been used.

16.1 Introduction

Chapter 15 discusses the functional programming paradigm, which is significantly different from the software development methodologies used with the imperative languages. In this chapter, we describe another different programming methodology. In this case, the approach is to express programs in a form of symbolic logic and use a logical inferencing process to produce results. Logic programs are declarative rather than procedural, which means that only the specifications of the desired results are stated rather than detailed procedures for producing them. Programs in logic programming languages are collections of facts and rules. Such a program is used by asking it questions, which it attempts to answer by consulting the facts and rules. “Consulting” is perhaps misleading here, for the process is far more complex than that word connotes. This approach to problem solving may sound like it addresses only a very narrow category of problems, but it is more flexible than might be thought.

Programming that uses a form of symbolic logic as a programming language is often called logic programming, and languages based on symbolic logic are called logic programming languages, or declarative languages. We have chosen to describe the logic programming language Prolog, because it is the only widely used logic language.

The syntax of logic programming languages is remarkably different from that of the imperative and functional languages. The semantics of logic programs also bears little resemblance to that of imperative-language programs. These observations should lead the reader to some curiosity about the nature of logic programming and declarative languages.

16.2 A Brief Introduction to Predicate Calculus

Before we can discuss logic programming, we must briefly investigate its basis, which is formal logic. This is not our first contact with formal logic in this book; it was used extensively in the axiomatic semantics described in Chapter 3.
A proposition can be thought of as a logical statement that may or may not be true. It consists of objects and the relationships among objects. Formal logic was developed to provide a method for describing propositions, with the goal of allowing those formally stated propositions to be checked for validity.

Symbolic logic can be used for the three basic needs of formal logic: to express propositions, to express the relationships between propositions, and to describe how new propositions can be inferred from other propositions that are assumed to be true.

There is a close relationship between formal logic and mathematics. In fact, much of mathematics can be thought of in terms of logic. The fundamental axioms of number and set theory are the initial set of propositions, which are assumed to be true. Theorems are the additional propositions that can be inferred from the initial set.

The particular form of symbolic logic that is used for logic programming is called first-order predicate calculus (though it is a bit imprecise, we will usually refer to it as predicate calculus). In the following subsections, we present a brief look at predicate calculus. Our goal is to lay the groundwork for a discussion of logic programming and the logic programming language Prolog.

16.2.1 Propositions

The objects in logic programming propositions are represented by simple terms, which are either constants or variables. A constant is a symbol that represents an object. A variable is a symbol that can represent different objects at different times, although in a sense that is far closer to mathematics than the variables in an imperative programming language.

The simplest propositions, which are called atomic propositions, consist of compound terms. A compound term is one element of a mathematical relation, written in a form that has the appearance of mathematical function notation. Recall from Chapter 15 that a mathematical function is a mapping, which can be represented either as an expression or as a table or list of tuples. Compound terms are elements of the tabular definition of a function.

A compound term is composed of two parts: a functor, which is the function symbol that names the relation, and an ordered list of parameters, which together represent an element of the relation. A compound term with a single parameter is a 1-tuple; one with two parameters is a 2-tuple, and so forth. For example, we might have the two propositions

\[
\text{man (jake)} \\
\text{like (bob, steak)}
\]

which state that \{jake\} is a 1-tuple in the relation named man, and that \{bob, steak\} is a 2-tuple in the relation named like. If we added the proposition

\[
\text{man (fred)}
\]
to the two previous propositions, then the relation man would have two distinct elements, \{jake\} and \{fred\}. All of the simple terms in these propositions—man, jake, like, bob, and steak—are constants. Note that these propositions have no intrinsic semantics. They mean whatever we want them to mean. For example, the second example may mean that bob likes steak, or that steak likes bob, or that bob is in some way similar to a steak.

Propositions can be stated in two modes: one in which the proposition is defined to be true, and one in which the truth of the proposition is something that is to be determined. In other words, propositions either can be facts or queries. The example propositions above could be either.

Compound propositions have two or more atomic propositions, which are connected by logical connectors, or operators, in the same way compound logic expressions are constructed in imperative languages. The names, symbols, and meanings of the predicate calculus logical connectors are as follows:

<table>
<thead>
<tr>
<th>Name</th>
<th>Symbol</th>
<th>Example</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>negation</td>
<td>(\neg)</td>
<td>(\neg a)</td>
<td>not (a)</td>
</tr>
<tr>
<td>conjunction</td>
<td>(\cap)</td>
<td>(a \cap b)</td>
<td>(a) and (b)</td>
</tr>
<tr>
<td>disjunction</td>
<td>(\cup)</td>
<td>(a \cup b)</td>
<td>(a) or (b)</td>
</tr>
<tr>
<td>equivalence</td>
<td>(\equiv)</td>
<td>(a \equiv b)</td>
<td>(a) is equivalent to (b)</td>
</tr>
<tr>
<td>implication</td>
<td>(\supset)</td>
<td>(a \supset b)</td>
<td>(a) implies (b)</td>
</tr>
<tr>
<td></td>
<td>(\subset)</td>
<td>(a \subset b)</td>
<td>(b) implies (a)</td>
</tr>
</tbody>
</table>

The following are examples of compound propositions:

\[a \cap b \supset c\]
\[a \cap \neg b \supset d\]

The \(\neg\) operator has the highest precedence. The operators \(\cap\), \(\cup\), and \(\equiv\) all have higher precedence than \(\supset\) and \(\subset\). So, the second example above is equivalent to

\[(a \cap (\neg b)) \supset d\]

Variables can appear in propositions but only when introduced by special symbols called quantifiers. Predicate calculus includes two quantifiers, as described below, where \(X\) is a variable and \(P\) is a proposition:

<table>
<thead>
<tr>
<th>Name</th>
<th>Example</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>universal</td>
<td>(\forall X.P)</td>
<td>For all (X), (P) is true.</td>
</tr>
<tr>
<td>existential</td>
<td>(\exists X.P)</td>
<td>There exists a value of (X) such that (P) is true.</td>
</tr>
</tbody>
</table>
The period between \( X \) and \( P \) simply separates the variable from the proposition. For example, consider the following:

\[
\forall X. \text{(woman}(X) \supset \text{human}(X))
\]
\[
\exists X. \text{(mother}(\text{mary}, X) \land \text{male}(X))
\]

The first of these propositions means that for any value of \( X \), if \( X \) is a woman, then \( X \) is a human. The second means that there exists a value of \( X \) such that \( \text{mary} \) is the mother of \( X \) and \( X \) is a male; in other words, \( \text{mary} \) has a son. The scope of the universal and existential quantifiers is the atomic propositions to which they are attached. This scope can be extended using parentheses, as in the two compound propositions just described. So, the universal and existential quantifiers have higher precedence than any of the operators.

### 16.2.2 Clausal Form

We are discussing predicate calculus because it is the basis for logic programming languages. As with other languages, logic languages are best in their simplest form, meaning that redundancy should be minimized.

One problem with predicate calculus as we have described it thus far is that there are too many different ways of stating propositions that have the same meaning; that is, there is a great deal of redundancy. This is not such a problem for logicians, but if predicate calculus is to be used in an automated (computerized) system, it is a serious problem. To simplify matters, a standard form for propositions is desirable. Clausal form, which is a relatively simple form of propositions, is one such standard form. All propositions can be expressed in clausal form. A proposition in clausal form has the following general syntax:

\[
B_1 \lor B_2 \lor \ldots \lor B_n \subset A_1 \land A_2 \land \ldots \land A_m
\]

in which the \( A \)'s and \( B \)'s are terms. The meaning of this clausal form proposition is as follows: If all of the \( A \)'s are true, then at least one \( B \) is true. The primary characteristics of clausal form propositions are the following: Existential quantifiers are not required; universal quantifiers are implicit in the use of variables in the atomic propositions; and no operators other than conjunction and disjunction are required. Also, conjunction and disjunction need appear only in the order shown in the general clausal form: disjunction on the left side and conjunction on the right side. All predicate calculus propositions can be algorithmically converted to clausal form. Nilsson (1971) gives proof that this can be done, as well as a simple conversion algorithm for doing it.

The right side of a clausal form proposition is called the **antecedent**. The left side is called the **consequent** because it is the consequence of the truth of the antecedent. As examples of clausal form propositions, consider the following:

\[
\text{likes}(\text{bob, trout}) \subset \text{likes}(\text{bob, fish}) \land \text{fish}(\text{trout})
\]
father (louis, al) ∪ father (louis, violet) ⊆ father (al, bob) ∩ mother (violet, bob) ∩ grandfather (louis, bob)

The English version of the first of these states that if bob likes fish and a trout is a fish, then bob likes trout. The second states that if al is bob’s father and violet is bob’s mother and louis is bob’s grandfather, then louis is either al’s father or violet’s father.

16.3 Predicate Calculus and Proving Theorems

Predicate calculus provides a method of expressing collections of propositions. One use of collections of propositions is to determine whether any interesting or useful facts can be inferred from them. This is exactly analogous to the work of mathematicians, who strive to discover new theorems that can be inferred from known axioms and theorems.

The early days of computer science (the 1950s and early 1960s) saw a great deal of interest in automating the theorem-proving process. One of the most significant breakthroughs in automatic theorem proving was the discovery of the resolution principle by Alan Robinson (1965) at Syracuse University.

Resolution is an inference rule that allows inferred propositions to be computed from given propositions, thus providing a method with potential application to automatic theorem proving. Resolution was devised to be applied to propositions in clausal form. The concept of resolution is the following: Suppose there are two propositions with the forms

\[ P_1 \subseteq P_2 \]
\[ Q_1 \subseteq Q_2 \]

Their meaning is that \( P_2 \) implies \( P_1 \) and \( Q_2 \) implies \( Q_1 \). Furthermore, suppose that \( P_1 \) is identical to \( Q_2 \), so that we could rename \( P_1 \) and \( Q_2 \) as \( T \). Then, we could rewrite the two propositions as

\[ T \subseteq P_2 \]
\[ Q_1 \subseteq T \]

Now, because \( P_2 \) implies \( T \) and \( T \) implies \( Q_1 \), it is logically obvious that \( P_2 \) implies \( Q_1 \), which we could write as

\[ Q_1 \subseteq P_2 \]

The process of inferring this proposition from the original two propositions is resolution.

As another example, consider the two propositions:

\[ \text{older} (\text{joanne, jake}) \subseteq \text{mother} (\text{joanne, jake}) \]
\[ \text{wiser} (\text{joanne, jake}) \subseteq \text{older} (\text{joanne, jake}) \]

From these propositions, the following proposition can be constructed using resolution:
wiser (joanne, jake) ⊆ mother (joanne, jake)

The mechanics of this resolution construction are simple: The terms of the left sides of the two clausal propositions are OR’d together to make the left side of the new proposition. Then the right sides of the two clausal propositions are AND’d together to get the right side of the new proposition. Next, any term that appears on both sides of the new proposition is removed from both sides. The process is exactly the same when the propositions have multiple terms on either or both sides. The left side of the new inferred proposition initially contains all of the terms of the left sides of the two given propositions. The new right side is similarly constructed. Then the term that appears on both sides of the new proposition is removed. For example, if we have

father (bob, jake) ∪ mother (bob, jake) ⊆ parent (bob, jake)
grandfather (bob, fred) ⊆ father (bob, jake) ∩ father (jake, fred)

resolution says that

mother (bob, jake) ∪ grandfather (bob, fred) ⊆
parent (bob, jake) ∩ father (jake, fred)

which has all but one of the atomic propositions of both of the original propositions. The one atomic proposition that allowed the operation father (bob, jake) in the left side of the first and in the right side of the second is left out. In English, we would say

if: bob is the parent of jake implies that bob is either the father or mother of jake
and: bob is the father of jake and jake is the father of fred implies that bob is the grandfather of fred
then: if bob is the parent of jake and jake is the father of fred then: either bob is jake’s mother or bob is fred’s grandfather

Resolution is actually more complex than these simple examples illustrate. In particular, the presence of variables in propositions requires resolution to find values for those variables that allow the matching process to succeed. This process of determining useful values for variables is called unification. The temporary assigning of values to variables to allow unification is called instantiation.

It is common for the resolution process to instantiate a variable with a value, fail to complete the required matching, and then be required to backtrack and instantiate the variable with a different value. We will discuss unification and backtracking more extensively in the context of Prolog.

A critically important property of resolution is its ability to detect any inconsistency in a given set of propositions. This is based on the formal property of resolution called refutation complete. What this means is that given a set of inconsistent propositions, resolution can prove them to be inconsistent. This allows resolution to be used to prove theorems, which can be done as
follows: We can envision a theorem proof in terms of predicate calculus as a given set of pertinent propositions, with the negation of the theorem itself stated as a new proposition. The theorem is negated so that resolution can be used to prove the theorem by finding an inconsistency. This is proof by contradiction, a frequently used approach to proving theorems in mathematics. Typically, the original propositions are called the hypotheses, and the negation of the theorem is called the goal.

Theoretically, this process is valid and useful. The time required for resolution, however, can be a problem. Although resolution is a finite process when the set of propositions is finite, the time required to find an inconsistency in a large database of propositions may be huge.

Theorem proving is the basis for logic programming. Much of what is computed can be couched in the form of a list of given facts and relationships as hypotheses, and a goal to be inferred from the hypotheses, using resolution. Resolution on a hypotheses and a goal that are general propositions, even if they are in clausal form, is often not practical. Although it may be possible to prove a theorem using clausal form propositions, it may not happen in a reasonable amount of time. One way to simplify the resolution process is to restrict the form of the propositions. One useful restriction is to require the propositions to be Horn clauses. Horn clauses only can be in one of two forms: They have either a single atomic proposition on the left side or an empty left side. The left side of a clausal form proposition is sometimes called the head, and Horn clauses with left sides are called headed Horn clauses. Headed Horn clauses are used to state relationships, such as

\[
\text{likes}(\text{bob}, \text{trout}) \subseteq \text{likes}(\text{bob}, \text{fish}) \cap \text{fish}(\text{trout})
\]

Horn clauses with empty left sides, which are often used to state facts, are called headless Horn clauses. For example,

\[
\text{father}(\text{bob}, \text{jake})
\]

Most, but not all, propositions can be stated as Horn clauses. The restriction to Horn clauses makes resolution a practical process for proving theorems.

16.4 An Overview of Logic Programming

Languages used for logic programming are called declarative languages, because programs written in them consist of declarations rather than assignments and control flow statements. These declarations are actually statements, or propositions, in symbolic logic.

One of the essential characteristics of logic programming languages is their semantics, which is called declarative semantics. The basic concept of this semantics is that there is a simple way to determine the meaning of each statement, and it does not depend on how the statement might be used to solve a

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1. Horn clauses are named after Alfred Horn (1951), who studied clauses in this form.
problem. Declarative semantics is considerably simpler than the semantics of the imperative languages. For example, the meaning of a given proposition in a logic programming language can be concisely determined from the statement itself. In an imperative language, the semantics of a simple assignment statement requires examination of local declarations, knowledge of the scoping rules of the language, and possibly even examination of programs in other files just to determine the types of the variables in the assignment statement. Then, assuming the expression of the assignment contains variables, the execution of the program prior to the assignment statement must be traced to determine the values of those variables. The resulting action of the statement, then, depends on its run-time context. Comparing this semantics with that of a proposition in a logic language, with no need to consider textual context or execution sequences, it is clear that declarative semantics is far simpler than the semantics of imperative languages. Thus, declarative semantics is often stated as one of the advantages that declarative languages have over imperative languages (Hogger, 1984, pp. 240–241).

Programming in both imperative and functional languages is primarily procedural, which means that the programmer knows what is to be accomplished by a program and instructs the computer on exactly how the computation is to be done. In other words, the computer is treated as a simple device that obeys orders. Everything that is computed must have every detail of that computation spelled out. Some believe that this is the essence of the difficulty of programming using imperative and functional languages.

Programming in a logic programming language is nonprocedural. Programs in such languages do not state exactly how a result is to be computed but rather describe the form of the result. The difference is that we assume the computer system can somehow determine how the result is to be computed. What is needed to provide this capability for logic programming languages is a concise means of supplying the computer with both the relevant information and a method of inference for computing desired results. Predicate calculus supplies the basic form of communication to the computer, and resolution provides the inference technique.

Sorting is commonly used to illustrate the difference between procedural and nonprocedural systems. In a language like Java, sorting is done by explaining in a Java program all of the details of some sorting algorithm to a computer that has a Java compiler. The computer, after translating the Java program into machine code or some interpretive intermediate code, follows the instructions and produces the sorted list.

In a nonprocedural language, it is necessary only to describe the characteristics of the sorted list: It is some permutation of the given list such that for each pair of adjacent elements, a given relationship holds between the two elements. To state this formally, suppose the list to be sorted is in an array named list, has a subscript range 1 . . . n. The concept of sorting the elements of the given list, named old_list, and placing them in a separate array, named new_list, can then be expressed as follows:

\[
\text{sort} (\text{old_list}, \text{new_list}) \subseteq \text{permute} (\text{old_list}, \text{new_list}) \cap \text{sorted} (\text{new_list})
\]  
\[
\text{sorted} (\text{list}) \subseteq \forall j \text{ such that } 11 \leq j < n, \text{list}(j) \leq \text{list}(j + 1)
\]
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where permute is a predicate that returns true if its second parameter array is a permutation of its first parameter array.

From this description, the nonprocedural language system could produce the sorted list. That makes nonprocedural programming sound like the mere production of concise software requirements specifications, which is a fair assessment. Unfortunately, however, it is not that simple. Logic programs that use only resolution face serious problems of execution efficiency. In our example of sorting, if the list is long, the number of permutations is huge, and they must be generated and tested, one by one, until the one that is in order is found—a very lengthy process. Of course, one must consider the possibility that the best form of a logic language may not yet have been determined, and good methods of creating programs in logic programming languages for large problems have not yet been developed.

16.5 The Origins of Prolog

As was stated in Chapter 2, Alain Colmerauer and Phillippe Roussel at the University of Aix-Marseille, with some assistance from Robert Kowalski at the University of Edinburgh, developed the fundamental design of Prolog. Colmerauer and Roussel were interested in natural-language processing; Kowalski was interested in automated theorem proving. The collaboration between the University of Aix-Marseille and the University of Edinburgh continued until the mid-1970s. Since then, research on the development and use of the language has progressed independently at those two locations, resulting in, among other things, two syntactically different dialects of Prolog.

The development of Prolog and other research efforts in logic programming received limited attention outside of Edinburgh and Marseille until the announcement in 1981 that the Japanese government was launching a large research project called the Fifth Generation Computing Systems (FGCS; Fuchi, 1981; Moto-oka, 1981). One of the primary objectives of the project was to develop intelligent machines, and Prolog was chosen as the basis for this effort. The announcement of FGCS aroused a sudden strong interest in artificial intelligence and logic programming in researchers and the governments of the United States and several European countries.

After a decade of effort, the FGCS project was quietly dropped. Despite the great assumed potential of logic programming and Prolog, little of great significance had been discovered. This led to the decline in the interest in and use of Prolog, although it still has its proponents.

16.6 The Basic Elements of Prolog

There are now a number of different dialects of Prolog. These can be grouped into several categories: those that grew from the Marseille group, those that came from the Edinburgh group, and some dialects that have been developed
for microcomputers, such as micro-Prolog, which is described by Clark and McCabe (1984). The syntactic forms of these are somewhat different. Rather than attempt to describe the syntax of several dialects of Prolog or some hybrid of them, we have chosen one particular, widely available dialect, which is the one developed at Edinburgh. This form of the language is sometimes called Edinburgh syntax. Its first implementation was on a DEC System-10 (Warren et al., 1979). Prolog implementations are available for virtually all popular computer platforms, for example, from the Free Software Organization (http://www.gnu.org).

16.6.1 Terms

As with programs in other languages, Prolog programs consist of collections of statements. There are only a few kinds of statements in Prolog, but they can be complex. All Prolog statements, as well as Prolog data, are constructed from terms.

A Prolog term is a constant, a variable, or a structure. A constant is either an atom or an integer. Atoms are the symbolic values of Prolog and are similar to their counterparts in LISP. In particular, an atom is either a string of letters, digits, and underscores that begins with a lowercase letter or a string of any printable ASCII characters delimited by apostrophes.

A variable is any string of letters, digits, and underscores that begins with an uppercase letter or an underscore (_). Variables are not bound to types by declarations. The binding of a value, and thus a type, to a variable is called an instantiation. Instantiation occurs only in the resolution process. A variable that has not been assigned a value is called uninstantiated. Instantiations last only as long as it takes to satisfy one complete goal, which involves the proof or disproof of one proposition. Prolog variables are only distant relatives, in terms of both semantics and use, to the variables in the imperative languages.

The last kind of term is called a structure. Structures represent the atomic propositions of predicate calculus, and their general form is the same:

\[ \text{functor} (\text{parameter list}) \]

The functor is any atom and is used to identify the structure. The parameter list can be any list of atoms, variables, or other structures. As discussed at length in the following subsection, structures are the means of specifying facts in Prolog. They can also be thought of as objects, in which case they allow facts to be stated in terms of several related atoms. In this sense, structures are relations, for they state relationships among terms. A structure is also a predicate when its context specifies it to be a query (question).

16.6.2 Fact Statements

Our discussion of Prolog statements begins with those statements used to construct the hypotheses, or database of assumed information—the statements from which new information can be inferred.
Prolog has two basic statement forms; these correspond to the headless and headed Horn clauses of predicate calculus. The simplest form of headless Horn clause in Prolog is a single structure, which is interpreted as an unconditional assertion, or fact. Logically, facts are simply propositions that are assumed to be true.

The following examples illustrate the kinds of facts one can have in a Prolog program. Notice that every Prolog statement is terminated by a period.

```
female(shelley).
male(bill).
female(mary).
male(jake).
father(bill, jake).
father(bill, shelley).
mother(mary, jake).
mother(mary, shelley).
```

These simple structures state certain facts about jake, shelley, bill, and mary. For example, the first states that shelley is a female. The last four connect their two parameters with a relationship that is named in the functor atom; for example, the fifth proposition might be interpreted to mean that bill is the father of jake. Note that these Prolog propositions, like those of predicate calculus, have no intrinsic semantics. They mean whatever the programmer wants them to mean. For example, the proposition

```
father(bill, jake).
```

could mean bill and jake have the same father or that jake is the father of bill. The most common and straightforward meaning, however, might be that bill is the father of jake.

### 16.6.3 Rule Statements

The other basic form of Prolog statement for constructing the database corresponds to a headed Horn clause. This form can be related to a known theorem in mathematics from which a conclusion can be drawn if the set of given conditions is satisfied. The right side is the antecedent, or if part, and the left side is the consequent, or then part. If the antecedent of a Prolog statement is true, then the consequent of the statement must also be true. Because they are Horn clauses, the consequent of a Prolog statement is a single term, while the antecedent can be either a single term or a conjunction.

**Conjunctions** contain multiple terms that are separated by logical AND operations. In Prolog, the AND operation is implied. The structures that specify atomic propositions in a conjunction are separated by commas, so one could consider the commas to be AND operators. As an example of a conjunction, consider the following:
female(shelley), child(shelley).

The general form of the Prolog headed Horn clause statement is

\[ \text{consequence} :- \text{antecedent_expression}. \]

It is read as follows: “consequence can be concluded if the antecedent expression is true or can be made to be true by some instantiation of its variables.” For example,

\[ \text{ancestor(mary, shelley)} :- \text{mother(mary, shelley)}. \]

states that if mary is the mother of shelley, then mary is an ancestor of shelley. Headed Horn clauses are called rules, because they state rules of implication between propositions.

As with clausal form propositions in predicate calculus, Prolog statements can use variables to generalize their meaning. Recall that variables in clausal form provide a kind of implied universal quantifier. The following demonstrates the use of variables in Prolog statements:

\[
\begin{align*}
\text{parent}(X, Y) & :- \text{mother}(X, Y). \\
\text{parent}(X, Y) & :- \text{father}(X, Y). \\
\text{grandparent}(X, Z) & :- \text{parent}(X, Y) , \text{parent}(Y, Z).
\end{align*}
\]

These statements give rules of implication among some variables, or universal objects. In this case, the universal objects are X, Y, and Z. The first rule states that if there are instantiations of X and Y such that mother(X, Y) is true, then for those same instantiations of X and Y, parent(X, Y) is true.

The = operator, which is an infix operator, succeeds if its two term operands are the same. For example, \( X = Y \). The not operator, which is a unary operator, reverses its operand, in the sense that it succeeds if its operand fails. For example, \( \text{not}(X = Y) \) succeeds if X is not equal to Y.

### 16.6.4 Goal Statements

So far, we have described the Prolog statements for logical propositions, which are used to describe both known facts and rules that describe logical relationships among facts. These statements are the basis for the theorem-proving model. The theorem is in the form of a proposition that we want the system to either prove or disprove. In Prolog, these propositions are called goals, or queries. The syntactic form of Prolog goal statements is identical to that of headless Horn clauses. For example, we could have

\[ \text{man(fred)}. \]

to which the system will respond either yes or no. The answer yes means that the system has proved the goal was true under the given database of facts and
relationships. The answer no means that either the goal was determined to be false or the system was simply unable to prove it.

Conjunctive propositions and propositions with variables are also legal goals. When variables are present, the system not only asserts the validity of the goal but also identifies the instantiations of the variables that make the goal true. For example,

\texttt{father}(X, \texttt{mike}).

can be asked. The system will then attempt, through unification, to find an instantiation of \( X \) that results in a true value for the goal.

Because goal statements and some nongoal statements have the same form (headless Horn clauses), a Prolog implementation must have some means of distinguishing between the two. Interactive Prolog implementations do this by simply having two modes, indicated by different interactive prompts: one for entering fact and rule statements and one for entering goals. The user can change the mode at any time.

### 16.6.5 The Inferencing Process of Prolog

This section examines Prolog resolution. Efficient use of Prolog requires that the programmer know precisely what the Prolog system does with his or her program.

Queries are called goals. When a goal is a compound proposition, each of the facts (structures) is called a subgoal. To prove that a goal is true, the inferencing process must find a chain of inference rules and/or facts in the database that connect the goal to one or more facts in the database. For example, if \( Q \) is the goal, then either \( Q \) must be found as a fact in the database or the inferencing process must find a fact \( P_1 \) and a sequence of propositions \( P_2, P_3, \ldots, P_n \) such that

\[
P_2 :\neg P_1
P_3 :\neg P_2
\ldots
Q :\neg P_n
\]

Of course, the process can be and often is complicated by rules with compound right sides and rules with variables. The process of finding the \( P_s \), when they exist, is basically a comparison, or matching, of terms with each other.

Because the process of proving a subgoal is done through a proposition-matching process, it is sometimes called matching. In some cases, proving a subgoal is called satisfying that subgoal.

Consider the following query:

\texttt{man} (bob).

This goal statement is the simplest kind. It is relatively easy for resolution to determine whether it is true or false: The pattern of this goal is compared with the facts and rules in the database. If the database includes the fact
The proof is trivial. If, however, the database contains the following fact and inference rule,

\[ \text{father}(\text{bob}). \]
\[ \text{man}(X) :\text{father}(X). \]

Prolog would be required to find these two statements and use them to infer the truth of the goal. This would necessitate unification to instantiate \( X \) temporarily to \( \text{bob} \).

Now consider the goal

\[ \text{man}(X). \]

In this case, Prolog must match the goal against the propositions in the database. The first proposition that it finds that has the form of the goal, with any object as its parameter, will cause \( X \) to be instantiated with that object’s value. \( X \) is then displayed as the result. If there is no proposition having the form of the goal, the system indicates, by saying \text{no}, that the goal cannot be satisfied.

There are two opposite approaches to attempting to match a given goal to a fact in the database. The system can begin with the facts and rules of the database and attempt to find a sequence of matches that lead to the goal. This approach is called \text{bottom-up resolution}, or \text{forward chaining}. The alternative is to begin with the goal and attempt to find a sequence of matching propositions that lead to some set of original facts in the database. This approach is called \text{top-down resolution}, or \text{backward chaining}. In general, backward chaining works well when there is a reasonably small set of candidate answers. The forward chaining approach is better when the number of possibly correct answers is large; in this situation, backward chaining would require a very large number of matches to get to an answer. Prolog implementations use backward chaining for resolution, presumably because its designers believed backward chaining was more suitable for a larger class of problems than forward chaining.

The following example illustrates the difference between forward and backward chaining. Consider the query:

\[ \text{man}(\text{bob}). \]

Assume the database contains

\[ \text{father}(\text{bob}). \]
\[ \text{man}(X) :\text{father}(X). \]

Forward chaining would search for and find the first proposition. The goal is then inferred by matching the first proposition with the right side of the second rule \( \text{father}(X) \) through instantiation of \( X \) to \( \text{bob} \) and then matching the left side of the second proposition to the goal. Backward chaining would first
match the goal with the left side of the second proposition \( \text{man}(X) \) through the instantiation of \( X \) to \( \text{bob} \). As its last step, it would match the right side of the second proposition (now \( \text{father}(\text{bob}) \)) with the first proposition.

The next design question arises whenever the goal has more than one structure, as in our example. The question then is whether the solution search is done depth first or breadth first. A \textit{depth-first} search finds a complete sequence of propositions—a proof—for the first subgoal before working on the others. A \textit{breadth-first} search works on all subgoals of a given goal in parallel. Prolog's designers chose the depth-first approach primarily because it can be done with fewer computer resources. The breadth-first approach is a parallel search that can require a large amount of memory.

The last feature of Prolog's resolution mechanism that must be discussed is backtracking. When a goal with multiple subgoals is being processed and the system fails to show the truth of one of the subgoals, the system abandons the subgoal it cannot prove. It then reconsiders the previous subgoal, if there is one, and attempts to find an alternative solution to it. This backing up in the goal to the reconsideration of a previously proven subgoal is called \textit{backtracking}. A new solution is found by beginning the search where the previous search for that subgoal stopped. Multiple solutions to a subgoal result from different instantiations of its variables. Backtracking can require a great deal of time and space because it may have to find all possible proofs to every subgoal. These subgoal proofs may not be organized to minimize the time required to find the one that will result in the final complete proof, which exacerbates the problem.

To solidify your understanding of backtracking, consider the following example. Assume that there is a set of facts and rules in the database and that Prolog has been presented with the following compound goal:

\[
\text{male}(X), \text{parent}(X, \text{shelley}).
\]

This goal asks whether there is an instantiation of \( X \) such that \( X \) is a \text{male} and \( X \) is a \text{parent} of \( \text{shelley} \). As its first step, Prolog finds the first fact in the database with \text{male} as its functor. It then instantiates \( X \) to the parameter of the found fact, say \( \text{mike} \). Then, it attempts to prove that \( \text{parent}(\text{mike}, \text{shelley}) \) is true. If it fails, it backtracks to the first subgoal, \( \text{male}(X) \), and attempts to resatisfy it with some alternative instantiation of \( X \). The resolution process may have to find every \text{male} in the database before it finds the one that is a \text{parent} of \( \text{shelley} \). It definitely must find all \text{males} to prove that the goal cannot be satisfied. Note that our example goal might be processed more efficiently if the order of the two subgoals were reversed. Then, only after resolution had found a \text{parent} of \( \text{shelley} \) would it try to match that person with the \text{male} subgoal. This is more efficient if \( \text{shelley} \) has fewer parents than there are \text{males} in the database, which seems like a reasonable assumption. Section 16.7.1 discusses a method of limiting the backtracking done by a Prolog system.

Database searches in Prolog always proceed in the direction of first to last.
The following two subsections describe Prolog examples that further illustrate the resolution process.

### 16.6.6 Simple Arithmetic

Prolog supports integer variables and integer arithmetic. Originally, the arithmetic operators were functors, so that the sum of 7 and the variable \( x \) was formed with

\[ +(7, x) \]

Prolog now allows a more abbreviated syntax for arithmetic with the `is` operator. This operator takes an arithmetic expression as its right operand and a variable as its left operand. All variables in the expression must already be instantiated, but the left-side variable cannot be previously instantiated. For example, in

\[ A \text{ is } B / 17 + C. \]

if \( B \) and \( C \) are instantiated but \( A \) is not, then this clause will cause \( A \) to be instantiated with the value of the expression. When this happens, the clause is satisfied. If either \( B \) or \( C \) is not instantiated or \( A \) is instantiated, the clause is not satisfied and no instantiation of \( A \) can take place. The semantics of an `is` proposition is considerably different from that of an assignment statement in an imperative language. This difference can lead to an interesting scenario. Because the `is` operator makes the clause in which it appears look like an assignment statement, a beginning Prolog programmer may be tempted to write a statement such as

\[ \text{Sum is Sum + Number.} \]

which is never useful, or even legal, in Prolog. If \( \text{Sum} \) is not instantiated, the reference to it in the right side is undefined and the clause fails. If \( \text{Sum} \) is already instantiated, the clause fails, because the left operand cannot have a current instantiation when `is` is evaluated. In either case, the instantiation of \( \text{Sum} \) to the new value will not take place. (If the value of \( \text{Sum + Number} \) is required, it can be bound to some new name.)

Prolog does not have assignment statements in the same sense as imperative languages. They are simply not needed in most of the programming for which Prolog was designed. The usefulness of assignment statements in imperative languages often depends on the capability of the programmer to control the execution control flow of the code in which the assignment statement is embedded. Because this type of control is not always possible in Prolog, such statements are far less useful.

As a simple example of the use of numeric computation in Prolog, consider the following problem: Suppose we know the average speeds of several
automobiles on a particular racetrack and the amount of time they are on the track. This basic information can be coded as facts, and the relationship between speed, time, and distance can be written as a rule, as in the following:

speed(ford, 100).
speed(chevy, 105).
speed(dodge, 95).
speed(volvo, 80).
time(ford, 20).
time(chevy, 21).
time(dodge, 24).
time(volvo, 24).
distance(X, Y) :- speed(X, Speed),
time(X, Time),
Y is Speed * Time.

Now, queries can request the distance traveled by a particular car. For example, the query

distance(chevy, Chevy_Distance).

instantiates Chevy_Distance with the value 2205. The first two clauses in the right side of the distance computation statement instantiate the variables Speed and Time with the corresponding values of the given automobile functor. After satisfying the goal, Prolog also displays the name Chevy_Distance and its value.

At this point it is instructive to take an operational look at how a Prolog system produces results. Prolog has a built-in structure named trace that displays the instantiations of values to variables at each step during the attempt to satisfy a given goal. trace is used to understand and debug Prolog programs. To understand trace, it is best to introduce a different model of the execution of Prolog programs, called the tracing model.

The tracing model describes Prolog execution in terms of four events: (1) call, which occurs at the beginning of an attempt to satisfy a goal, (2) exit, which occurs when a goal has been satisfied, (3) redo, which occurs when backtrack causes an attempt to resatisfy a goal, and (4) fail, which occurs when a goal fails. Call and exit can be related directly to the execution model of a subprogram in an imperative language if processes like distance are thought of as subprograms. The other two events are unique to logic programming systems. In the following trace example, a trace of the computation of the value for Chevy_Distance, the goal requires no redo or fail events:

trace.
distance(chevy, Chevy_Distance).

(1) 1 Call: distance(chevy, _0)?
(2) 2 Call: speed(chevy, _5)?
Symbols in the trace that begin with the underscore character (_ ) are internal variables used to store instantiated values. The first column of
the trace indicates the subgoal whose match is currently being attempted. For example, in the example trace, the first line with the indication (3)
is an attempt to instantiate the temporary variable _6 with a time value for
chevy, where time is the second term in the right side of the statement that
describes the computation of distance. The second column indicates the
call depth of the matching process. The third column indicates the current
action.

To illustrate backtracking, consider the following example database and
tagged compound goal:

likes(jake, chocolate).
likes(jake, apricots).
likes(darcie, licorice).
likes(darcie, apricots).

trace.
likes(jake, X), likes(darcie, X).

One can think about Prolog computations graphically as follows: Consider
each goal as a box with four ports—call, fail, exit, and redo. Control enters a
goal in the forward direction through its call port. Control can also enter a
goal from the reverse direction through its redo port. Control can also leave
a goal in two ways: If the goal succeeded, control leaves through the exit port;
if the goal failed, control leaves through the fail port. A model of the example
is shown in Figure 16.1. In this example, control flows through each subgoal
twice. The second subgoal fails the first time, which forces a return through redo to the first subgoal.

### 16.6.7 List Structures

So far, the only Prolog data structure we have discussed is the atomic proposition, which looks more like a function call than a data structure. Atomic propositions, which are also called structures, are actually a form of records. The other basic data structure supported is the list. Lists are sequences of any number of elements, where the elements can be atoms, atomic propositions, or any other terms, including other lists.

Prolog uses the syntax of ML and Haskell to specify lists. The list elements are separated by commas, and the entire list is delimited by square brackets, as in

\[
\text{[apple, prune, grape, kumquat]}
\]

The notation \([]\) is used to denote the empty list. Instead of having explicit functions for constructing and dismantling lists, Prolog simply uses a special notation. \([X \mid Y]\) denotes a list with head \(X\) and tail \(Y\), where head and tail correspond to \text{CAR} and \text{CDR} in LISP. This is similar to the notation used in ML and Haskell.

A list can be created with a simple structure, as in

\[
\text{new_list([apple, prune, grape, kumquat])}.
\]

which states that the constant list \([apple, prune, grape, kumquat]\) is a new element of the relation named \text{new_list} (a name we just made up). This statement does not bind the list to a variable named \text{new_list}; rather, it does the kind of thing that the proposition

\[
\text{likes(jake, X), likes(darcie, X)}
\]
male(jake)

does. That is, it states that [apple, prune, grape, kumquat] is a new element of new_list. Therefore, we could have a second proposition with a list argument, such as

new_list([apricot, peach, pear])

In query mode, one of the elements of new_list can be dismantled into head and tail with

new_list([New_List_Head | New_List_Tail]).

If new_list has been set to have the two elements as shown, this statement instantiates New_List_Head with the head of the first list element (in this case, apple) and New_List_Tail with the tail of the list (or [prune, grape, kumquat]). If this were part of a compound goal and backtracking forced a new evaluation of it, New_List_Head and New_List_Tail would be reinstantiated to apricot and [peach, pear], respectively, because [apricot, peach, pear] is the next element of new_list.

The | operator used to dismantle lists can also be used to create lists from given instantiated head and tail components, as in

[Element_1 | List_2]

If Element_1 has been instantiated with pickle and List_2 has been instantiated with [peanut, prune, popcorn], the sample notation will create, for this one reference, the list [pickle, peanut, prune, popcorn].

As stated previously, the list notation that includes the | symbol is universal: It can specify either a list construction or a list dismantling. Note further that the following are equivalent:

[apricot, peach, pear | []]
[apricot, peach | [pear]]
[apricot | [peach, pear]]

With lists, certain basic operations are often required, such as those found in LISP, ML, and Haskell. As an example of such operations in Prolog, we examine a definition of append, which is related to such a function in LISP. In this example, the differences and similarities between functional and declarative languages can be seen. We need not specify how Prolog is to construct a new list from the given lists; rather, we need specify only the characteristics of the new list in terms of the given lists.

In appearance, the Prolog definition of append is very similar to the ML version that appears in Chapter 15, and a kind of recursion in resolution is used in a similar way to produce the new list. In the case of Prolog, the recursion is caused and controlled by the resolution process. As with ML and Haskell,
a pattern-matching process is used to choose, based on the actual parameter, between two different definitions of the append process.

The first two parameters to the `append` operation in the following code are the two lists to be appended, and the third parameter is the resulting list:

```prolog
append([], List, List).
append([Head | List_1], List_2, [Head | List_3]) :-
    append(List_1, List_2, List_3).
```

The first proposition specifies that when the empty list is appended to any other list, that other list is the result. This statement corresponds to the recursion-terminating step of the ML `append` function. Note that the terminating proposition is placed before the recursion proposition. This is done because we know that Prolog will match the two propositions in order, starting with the first (because of its use of the depth-first order).

The second proposition specifies several characteristics of the new list. It corresponds to the recursion step in the ML function. The left-side predicate states that the first element of the new list is the same as the first element of the first given list, because they are both named `Head`. Whenever `Head` is instantiated to a value, all occurrences of `Head` in the goal are, in effect, simultaneously instantiated to that value. The right side of the second statement specifies that the tail of the first given list (List_1) has the second given list (List_2) appended to it to form the tail (List_3) of the resulting list.

One way to read the second statement of `append` is as follows: Appending the list `[Head | List_1]` to any list `List_2` produces the list `[Head | List_3]`, but only if the list `List_3` is formed by appending `List_1` to `List_2`. In LISP, this would be

```lisp
(CONS (CAR FIRST) (APPEND (CDR FIRST) SECOND))
```

In both the Prolog and LISP versions, the resulting list is not constructed until the recursion produces the terminating condition; in this case, the first list must become empty. Then, the resulting list is built using the `append` function itself; the elements taken from the first list are added, in reverse order, to the second list. The reversing is done by the unraveling of the recursion.

One fundamental difference between Prolog’s `append` and those of LISP and ML is that Prolog’s `append` is a predicate—it does not return a list, it returns yes or no. The new list is the value of its third parameter.

To illustrate how the `append` process progresses, consider the following traced example:

```prolog
trace.
append([bob, jo], [jake, darcie], Family).
```

(1) 1 Call: append([bob, jo], [jake, darcie], _10)?
(2) 2 Call: append([jo], [jake, darcie], _18)?
(3) 3 Call: append([], [jake, darcie], _25)?
(3) 3 Exit: append([], [jake, darcie], [jake, darcie])
(2) 2 Exit: append([jo], [jake, darcie], [jo, jake, darcie])
(1) 1 Exit: append([bob, jo], [jake, darcie], [bob, jo, jake, darcie])

Family = [bob, jo, jake, darcie]
yes

The first two calls, which represent subgoals, have List_1 nonempty, so they create the recursive calls from the right side of the second statement. The left side of the second statement effectively specifies the arguments for the recursive calls, or goals, thus dismantling the first list one element per step. When the first list becomes empty, in a call, or subgoal, the current instance of the right side of the second statement succeeds by matching the first statement. The effect of this is to return as the third parameter the value of the empty list appended to the second original parameter list. On successive exits, which represent successful matches, the elements that were removed from the first list are appended to the resulting list, Family. When the exit from the first goal is accomplished, the process is complete, and the resulting list is displayed.

Another difference between Prolog’s append and those of LISP and ML is that Prolog’s append is more flexible than that of those languages. For example, in Prolog we can use append to determine what two lists can be appended to get [a, b, c] with

append(X, Y, [a, b, c]).

This results in the following:

X = []
Y = [a, b, c]

If we type a semicolon at this output we get the alternative result:

X = [a]
Y = [b, c]

Continuing, we get the following:

X = [a, b]
Y = [c];
X = [a, b, c]
Y = []
The append predicate can also be used to create other list operations, such as the following, whose effect we invite the reader to determine. Note that list_op_2 is meant to be used by providing a list as its first parameter and a variable as its second, and the result of list_op_2 is the value to which the second parameter is instantiated.

list_op_2([], []).  
list_op_2([Head | Tail], List) :-  
list_op_2(Tail, Result), append(Result, [Head], List).

As you may have been able to determine, list_op_2 causes the Prolog system to instantiate its second parameter with a list that has the elements of the list of the first parameter, but in reverse order. For example, ([apple, orange, grape], Q) instantiates Q with the list [grape, orange, apple].

Once again, although the LISP and Prolog languages are fundamentally different, similar operations can use similar approaches. In the case of the reverse operation, both the Prolog's list_op_2 and LISP's reverse function include the recursion-terminating condition, along with the basic process of appending the reversal of the CDR or tail of the list to the CAR or head of the list to create the result list.

The following is a trace of this process, now named reverse:

trace. 
reverse([a, b, c], Q).

(1) 1 Call: reverse([a, b, c], _6)?  
(2) 2 Call: reverse([b, c], _65636)?  
(3) 3 Call: reverse([c], _65646)?  
(4) 4 Call: reverse([], _65656)?  
(4) 4 Exit: reverse([], [])  
(5) 4 Call: append([], [c], _65646)?  
(5) 4 Exit: append([], [c], [c])  
(3) 3 Exit: reverse([c], [c])  
(6) 3 Call: append([c], [b], _65636)?  
(7) 4 Call: append([], [b], _25)?  
(7) 4 Exit: append([], [b], [b])  
(6) 3 Exit: append([c], [b], [c, b])  
(2) 2 Exit: reverse([b, c], [c, b])  
(8) 2 Call: append([c, b], [a], _6)?  
(9) 3 Call: append([b], [a], _32)?  
(10) 4 Call: append([], [a], _39)?  
(10) 4 Exit: append([], [a], [a])  
(9) 3 Exit: append([b], [a], [b, a])  
(8) 2 Exit: append([c, b], [a], [c, b, a])  
(1) 1 Exit: reverse([a, b, c], [c, b, a])

Q = [c, b, a]
Suppose we need to be able to determine whether a given symbol is in a given list. A straightforward Prolog description of this is

```
member(Element, [Element | _]).
member(Element, [_ | List]) :- member(Element, List).
```

The underscore indicates an “anonymous” variable; it is used to mean that we do not care what instantiation it might get from unification. The first statement in the previous example succeeds if `Element` is the head of the list, either initially or after several recursions through the second statement. The second statement succeeds if `Element` is in the tail of the list. Consider the following traced examples:

```
trace.
member(a, [b, c, d]).
(1) 1 Call: member(a, [b, c, d])?
(2) 2 Call: member(a, [c, d])?
(3) 3 Call: member(a, [d])?
(4) 4 Call: member(a, [])?
(4) 4 Fail: member(a, [])
(3) 3 Fail: member(a, [d])
(2) 2 Fail: member(a, [c, d])
(1) 1 Fail: member(a, [b, c, d])
no

member(a, [b, a, c]).
(1) 1 Call: member(a, [b, a, c])?
(2) 2 Call: member(a, [a, c])?
(2) 2 Exit: member(a, [a, c])
(1) 1 Exit: member(a, [b, a, c])
yes
```

**16.7 Deficiencies of Prolog**

Although Prolog is a useful tool, it is neither a pure nor a perfect logic programming language. This section describes some of the problems with Prolog.

**16.7.1 Resolution Order Control**

Prolog, for reasons of efficiency, allows the user to control the ordering of pattern matching during resolution. In a pure logic programming environment, the order of attempted matches that take place during resolution is nondeterministic, and all matches could be attempted concurrently. However, because Prolog always matches in the same order, starting at the beginning of the database and at the left end of a given goal, the user can profoundly affect efficiency...
by ordering the database statements to optimize a particular application. For example, if the user knows that certain rules are much more likely to succeed than the others during a particular “execution,” then the program can be made more efficient by placing those rules at the beginning of the database.

In addition to allowing the user to control database and subgoal ordering, Prolog, in another concession to efficiency, allows some explicit control of backtracking. This is done with the cut operator, which is specified by an exclamation point (!). The cut operator is actually a goal, not an operator. As a goal, it always succeeds immediately, but it cannot be resatisfied through backtracking. Thus, a side effect of the cut is that subgoals to its left in a compound goal also cannot be resatisfied through backtracking. For example, in the goal

\[ a, b, !, c, d. \]

if both \( a \) and \( b \) succeed but \( c \) fails, the whole goal fails. This goal would be used if it were known that whenever \( c \) fails, it is a waste of time to resatisfy \( b \) or \( a \).

The purpose of the cut then is to allow the user to make programs more efficient by telling the system when it should not attempt to resatisfy subgoals that presumably could not result in a complete proof.

As an example of the use of the cut operator, consider the member rules from Section 16.6.7, which are:

\[
\begin{align*}
\text{member}(\text{Element}, [\text{Element} | _]) & . \\
\text{member}(\text{Element}, [\_ | \text{List}]) & : - \text{member}(\text{Element}, \text{List}).
\end{align*}
\]

If the list argument to \text{member} represents a set, then it can be satisfied only once (sets contain no duplicate elements). Therefore, if \text{member} is used as a subgoal in a multiple subgoal goal statement, there can be a problem. The problem is that if \text{member} succeeds but the next subgoal fails, backtracking will attempt to resatisfy \text{member} by continuing a prior match. But because the list argument to \text{member} has only one copy of the element to begin with, \text{member} cannot possibly succeed again, which eventually causes the whole goal to fail, in spite of any additional attempts to resatisfy \text{member}. For example, consider the goal:

\[
\text{dem_candidate}(X) : - \text{member}(X, \text{democrats}), \text{tests}(X).
\]

This goal determines whether a given person is a democrat and is a good candidate to run for a particular position. The \text{tests} subgoal checks a variety of characteristics of the given democrat to determine the suitability of the person for the position. If the set of democrats has no duplicates, then we do not want to back up to the \text{member} subgoal if the \text{tests} subgoal fails, because \text{member} will search all of the other democrats but fail, because there are no duplicates. The second attempt of \text{member} subgoal will be a waste of computation time. The solution to this inefficiency is to add a right side to the first statement of the \text{member} definition, with the cut operator as the sole element, as in
member(Element, [Element | _]) :- !.

Backtracking will not attempt to resatisfy \texttt{member} but instead will cause the entire subgoal to fail.

Cut is particularly useful in a programming strategy in Prolog called \texttt{generate and test}. In programs that use the generate-and-test strategy, the goal consists of subgoals that generate potential solutions, which are then checked by later “test” subgoals. Rejected solutions require backtracking to “generator” subgoals, which generate new potential solutions. As an example of a generate-and-test program, consider the following, which appears in Clocksin and Mellish (2003):

\begin{verbatim}
divide(N1, N2, Result) :- is_integer(Result),
    Product1 is Result * N2,
    Product2 is (Result + 1) * N2,
    Product1 =< N1, Product2 > N1, !.
\end{verbatim}

This program performs integer division, using addition and multiplication. Because most Prolog systems provide division as an operator, this program actually is not useful, other than to illustrate a simple generate-and-test program.

The predicate \texttt{is_integer} succeeds as long as its parameter can be instantiated to some nonnegative integer. If its argument is not instantiated, \texttt{is_integer} instantiates it to the value 0. If the argument is instantiated to an integer, \texttt{is_integer} instantiates it to the next larger integer value.

So, in \texttt{divide}, \texttt{is_integer} is the generator subgoal. It generates elements of the sequence 0, 1, 2, \ldots, one each time it is satisfied. All of the other subgoals are the testing subgoals—they check to determine whether the value produced by \texttt{is_integer} is, in fact, the quotient of the first two parameters, \texttt{N1} and \texttt{N2}. The purpose of the cut as the last subgoal is simple: It prevents \texttt{divide} from ever trying to find an alternative solution once it has found the solution. Although \texttt{is_integer} can generate a huge number of candidates, only one is the solution, so the cut here prevents useless attempts to produce secondary solutions.

Use of the cut operator has been compared to the use of the goto in imperative languages (van Emden, 1980). Although it is sometimes needed, it is possible to abuse it. Indeed, it is sometimes used to make logic programs have a control flow that is inspired by imperative programming styles.

The ability to tamper with control flow in a Prolog program is a deficiency, because it is directly detrimental to one of the important advantages of logic programming—that programs do not specify how solutions are to be found. Rather, they simply specify what the solution should look like. This design makes programs easier to write and easier to read. They are not cluttered with the details of how the solutions are to be determined and, in particular, the precise order in which the computations are done to produce the solution. So, while logic programming requires no control flow directions, Prolog programs frequently use them, mostly for the sake of efficiency.
16.7.2 The Closed-World Assumption

The nature of Prolog's resolution sometimes creates misleading results. The only truths, as far as Prolog is concerned, are those that can be proved using its database. It has no knowledge of the world other than its database. When the system receives a query and the database does not have information to prove the query absolutely, the query is assumed to be false. Prolog can prove that a given goal is true, but it cannot prove that a given goal is false. It simply assumes that, because it cannot prove a goal true, the goal must be false. In essence, Prolog is a true/fail system, rather than a true/false system.

Actually, the closed-world assumption should not be at all foreign to you—our judicial system operates the same way. Suspects are innocent until proven guilty. They need not be proven innocent. If a trial cannot prove a person guilty, he or she is considered innocent.

The problem of the closed-world assumption is related to the negation problem, which is discussed in the following subsection.

16.7.3 The Negation Problem

Another problem with Prolog is its difficulty with negation. Consider the following database of two facts and a relationship:

\[
\begin{align*}
\text{parent(bill, jake).} \\
\text{parent(bill, shelley).} \\
\text{sibling(X, Y) :- (parent(M, X), parent(M, Y).}
\end{align*}
\]

Now, suppose we typed the query

\[
\text{sibling(X, Y).}
\]

Prolog will respond with

\[
\begin{align*}
X &= jake \\
Y &= jake
\end{align*}
\]

Thus, Prolog “thinks” jake is a sibling of himself. This happens because the system first instantiates \(M\) with bill and \(X\) with jake to make the first subgoal, \(\text{parent(M, X)}\), true. It then starts at the beginning of the database again to match the second subgoal, \(\text{parent(M, Y)}\), and arrives at the instantiations of \(M\) with bill and \(Y\) with jake. Because the two subgoals are satisfied independently, with both matchings starting at the database’s beginning, the shown response appears. To avoid this result, \(X\) must be specified to be a sibling of \(Y\) only if they have the same parents and they are not the same. Unfortunately, stating that they are not equal is not straightforward in Prolog, as we will discuss. The most exacting method would require adding a fact for every pair of atoms, stating that they were not the same. This can, of course, cause the database to become very large, for there is often far more negative information than
positive information. For example, most people have 364 more unbirthdays than they have birthdays.

A simple alternative solution is to state in the goal that \( x \) must not be the same as \( y \), as in

\[
\text{sibling}(X, Y) :- \text{parent}(M, X), \text{parent}(M, Y), \text{not}(X = Y).
\]

In other situations, the solution is not so simple.

The Prolog \textit{not} operator is satisfied in this case if resolution cannot satisfy the subgoal \( X = Y \). Therefore, if \textit{not} succeeds, it does not necessarily mean that \( X \) is not equal to \( Y \); rather, it means that resolution cannot prove from the database that \( X \) is the same as \( Y \). Thus, the Prolog \textit{not} operator is not equivalent to a logical NOT operator, in which NOT means that its operand is probably true. This nonequivalency can lead to a problem if we happen to have a goal of the form

\[
\text{not}(\text{not}(\text{some}\_\text{goal})).
\]

which would be equivalent to

\[
\text{some}\_\text{goal}.
\]

if Prolog’s \textit{not} operator were a true logical NOT operator. In some cases, however, they are not the same. For example, consider again the \textit{member} rules:

\[
\begin{align*}
\text{member}(\text{Element}, [\text{Element} | \_]) & :- !. \\
\text{member}(\text{Element}, [\_ | \text{List}]) & :- \text{member}(\text{Element}, \text{List}).
\end{align*}
\]

To discover one of the elements of a given list, we could use the goal

\[
\text{member}(X, [\text{mary}, \text{fred}, \text{barb}]).
\]

which would cause \( X \) to be instantiated with \text{mary}, which would then be printed. But if we used

\[
\text{not}(\text{not}(\text{member}(X, [\text{mary}, \text{fred}, \text{barb}])))
\]

the following sequence of events would take place: First, the inner goal would succeed, instantiating \( X \) to \text{mary}. Then, Prolog would attempt to satisfy the next goal:

\[
\text{not}(\text{member}(X, [\text{mary}, \text{fred}, \text{barb}])).
\]

This statement would fail because \textit{member} succeeded. When this goal failed, \( X \) would be uninstantiated, because Prolog always uninstantiates all variables in all goals that fail. Next, Prolog would attempt to satisfy the outer \textit{not} goal,
which would succeed, because its argument had failed. Finally, the result, which is \( x \), would be printed. But \( x \) would not be currently instantiated, so the system would indicate that. Generally, uninstantiated variables are printed in the form of a string of digits preceded by an underscore. So, the fact that Prolog’s `not` is not equivalent to a logical NOT can be, at the very least, misleading.

The fundamental reason why logical NOT cannot be an integral part of Prolog is the form of the Horn clause:

\[
A \leftarrow B_1 \land B_2 \land \ldots \land B_n
\]

If all the \( B \) propositions are true, it can be concluded that \( A \) is true. But regardless of the truth or falseness of any or all of the \( B \)'s, it cannot be concluded that \( A \) is false. From positive logic, one can conclude only positive logic. Thus, the use of Horn clause form prevents any negative conclusions.

### 16.7.4 Intrinsic Limitations

A fundamental goal of logic programming, as stated in Section 16.4, is to provide nonprocedural programming; that is, a system by which programmers specify what a program is supposed to do but need not specify how that is to be accomplished. The example given there for sorting is rewritten here:

```prolog
sort (old_list, new_list) ⊆ permute (old_list, new_list) ∩ sorted (new_list)
sorted (list) ⊆ ∀j such that 1 ≤ j < n, list(j) ≤ list(j+1)
```

It is straightforward to write this in Prolog. For example, the sorted subgoal can be expressed as

```prolog
sorted ([]).
sorted ([x]).
sorted ([x, y | list]) :- x <= y, sorted ([y | list]).
```

The problem with this sort process is that it has no idea of how to sort, other than simply to enumerate all permutations of the given list until it happens to create the one that has the list in sorted order—a very slow process, indeed.

So far, no one has discovered a process by which the description of a sorted list can be transformed into some efficient algorithm for sorting. Resolution is capable of many interesting things, but certainly not this. Therefore, a Prolog program that sorts a list must specify the details of how that sorting can be done, as is the case in an imperative or functional language.

Do all of these problems mean that logic programming should be abandoned? Absolutely not! As it is, it is capable of dealing with many useful applications. Furthermore, it is based on an intriguing concept and is therefore interesting in and of itself. Finally, there is the possibility that new inferencing techniques will be developed that will allow a logic programming language system to efficiently deal with progressively larger classes of problems.
16.8 Applications of Logic Programming

In this section, we briefly describe a few of the larger classes of present and potential applications of logic programming in general and Prolog in particular.

16.8.1 Relational Database Management Systems

Relational database management systems (RDBMSs) store data in the form of tables. Queries on such databases are often stated in Structured Query Language (SQL). SQL is nonprocedural in the same sense that logic programming is nonprocedural. The user does not describe how to retrieve the answer; rather, he or she describes only the characteristics of the answer. The connection between logic programming and RDBMSs should be obvious. Simple tables of information can be described by Prolog structures, and relationships between tables can be conveniently and easily described by Prolog rules. The retrieval process is inherent in the resolution operation. The goal statements of Prolog provide the queries for the RDBMS. Logic programming is thus a natural match to the needs of implementing an RDBMS.

One of the advantages of using logic programming to implement an RDBMS is that only a single language is required. In a typical RDBMS, a database language includes statements for data definitions, data manipulation, and queries, all of which are embedded in a general-purpose programming language, such as COBOL. The general-purpose language is used for processing the data and input and output functions. All of these functions can be done in a logic programming language.

Another advantage of using logic programming to implement an RDBMS is that deductive capability is built in. Conventional RDBMSs cannot deduce anything from a database other than what is explicitly stored in them. They contain only facts, rather than facts and inference rules. The primary disadvantage of using logic programming for an RDBMS, compared with a conventional RDBMS, is that the logic programming implementation is slower. Logical inferences take longer than ordinary table look-up methods using imperative programming techniques.

16.8.2 Expert Systems

Expert systems are computer systems designed to emulate human expertise in some particular domain. They consist of a database of facts, an inferencing process, some heuristics about the domain, and some friendly human interface that makes the system appear much like an expert human consultant. In addition to their initial knowledge base, which is provided by a human expert, expert systems learn from the process of being used, so their databases must be capable of growing dynamically. Also, an expert system should include the capability of interrogating the user to get additional information when it determines that such information is needed.
One of the central problems for the designer of an expert system is dealing with the inevitable inconsistencies and incompleteness of the database. Logic programming appears to be well suited to deal with these problems. For example, default inference rules can help deal with the problem of incompleteness.

Prolog can and has been used to construct expert systems. It can easily fulfill the basic needs of expert systems, using resolution as the basis for query processing, using its ability to add facts and rules to provide the learning capability, and using its trace facility to inform the user of the “reasoning” behind a given result. Missing from Prolog is the automatic ability of the system to query the user for additional information when it is needed.

One of the most widely known uses of logic programming in expert systems is the expert system construction system known as APES, which is described in Sergot (1983) and Hammond (1983). The APES system includes a very flexible facility for gathering information from the user during expert system construction. It also includes a second interpreter for producing explanations to its answers to queries.

APES has been successfully used to produce several expert systems, including one for the rules of a government social benefits program and one for the British Nationality Act, which is the definitive source for rules of British citizenship.

### 16.8.3 Natural-Language Processing

Certain kinds of natural-language processing can be done with logic programming. In particular, natural-language interfaces to computer software systems, such as intelligent databases and other intelligent knowledge-based systems, can be conveniently done with logic programming. For describing language syntax, forms of logic programming have been found to be equivalent to context-free grammars. Proof procedures in logic programming systems have been found to be equivalent to certain parsing strategies. In fact, backward-chaining resolution can be used directly to parse sentences whose structures are described by context-free grammars. It has also been discovered that some kinds of semantics of natural languages can be made clear by modeling the languages with logic programming. In particular, research in logic-based semantics networks has shown that sets of sentences in natural languages can be expressed in clausal form (Deliyanni and Kowalski, 1979). Kowalski (1979) also discusses logic-based semantic networks.

### SUMMARY

Symbolic logic provides the basis for logic programming and logic programming languages. The approach of logic programming is to use as a database a collection of facts and rules that state relationships between facts and to use an automatic inferencing process to check the validity of new propositions,
assuming the facts and rules of the database are true. This approach is the one developed for automatic theorem proving.

Prolog is the most widely used logic programming language. The origins of logic programming lie in Robinson’s development of the resolution rule for logical inference. Prolog was developed primarily by Colmerauer and Roussel at Marseille, with some help from Kowalski at Edinburgh.

Logic programs are nonprocedural, which means that the characteristics of the solution are given but the process of getting the solution is not.

Prolog statements are facts, rules, or goals. Most are made up of structures, which are atomic propositions, and logic operators, although arithmetic expressions are also allowed.

Resolution is the primary activity of a Prolog interpreter. This process, which uses backtracking extensively, involves mainly pattern matching among propositions. When variables are involved, they can be instantiated to values to provide matches. This instantiation process is called unification.

There are a number of problems with the current state of logic programming. For reasons of efficiency, and even to avoid infinite loops, programmers must sometimes state control flow information in their programs. Also, there are the problems of the closed-world assumption and negation.

Logic programming has been used in a number of different areas, primarily in relational database systems, expert systems, and natural-language processing.

**BIBLIOGRAPHIC NOTES**

The Prolog language is described in several books. Edinburgh’s form of the language is covered in Clocksin and Mellish (2003). The microcomputer implementation is described in Clark and McCabe (1984).

Hogger (1991) is an excellent book on the general topic of logic programming. It is the source of the material in this chapter’s section on logic programming applications.

**REVIEW QUESTIONS**

1. What are the three primary uses of symbolic logic in formal logic?
2. What are the two parts of a compound term?
3. What are the two modes in which a proposition can be stated?
4. What is the general form of a proposition in clausal form?
5. What are antecedents? Consequences?
6. Give general (not rigorous) definitions of resolution and unification.
7. What are the forms of Horn clauses?
8. What is the basic concept of declarative semantics?
9. What does it mean for a language to be nonprocedural?
10. What are the three forms of a Prolog term?
11. What is an uninstantiated variable?
12. What are the syntactic forms and usage of fact and rule statements in Prolog?
13. What is a conjunction?
14. Explain the two approaches to matching goals to facts in a database.
15. Explain the difference between a depth-first and a breadth-first search when discussing how multiple goals are satisfied.
16. Explain how backtracking works in Prolog.
17. Explain what is wrong with the Prolog statement $K \text{ is } K + 1$.
18. What are the two ways a Prolog programmer can control the order of pattern matching during resolution?
19. Explain the generate-and-test programming strategy in Prolog.
20. Explain the closed-world assumption used by Prolog. Why is this a limitation?
21. Explain the negation problem with Prolog. Why is this a limitation?
22. Explain the connection between automatic theorem proving and Prolog’s inferencing process.
23. Explain the difference between procedural and nonprocedural languages.
24. Explain why Prolog systems must do backtracking.
25. What is the relationship between resolution and unification in Prolog?

**Problems Set**

1. Compare the concepts of procedural and nonprocedural systems.
2. Describe how a multiple-processor machine could be used to implement resolution. Could Prolog, as currently defined, use this method?
3. Write a Prolog description of your family tree (based only on facts), going back to your grandparents and including all descendants. Be sure to include all relationships.
4. Write a set of rules for family relationships, including all relationships from grandparents through two generations. Now add these to the facts of Problem 3, and eliminate as many of the facts as you can.
5. Write the following English conditional statements as Prolog-headed Horn clauses:
   a. If Fred is the father of Mike, then Fred is an ancestor of Mike.
   b. If Mike is the father of Joe and Mike is the father of Mary, then Mary is the sister of Joe.
   c. If Mike is the brother of Fred and Fred is the father of Mary, then Mike is the uncle of Mary.

6. Explain two ways in which the list-processing capabilities of Scheme and Prolog are similar.

7. In what way are the list-processing capabilities of Scheme and Prolog different?

8. Write a comparison of Prolog with ML, including two similarities and two differences.

9. From a book on Prolog, learn and write a description of an occur-check problem. Why does Prolog allow this problem to exist in its implementation?

10. Find a good source of information on intelligent database and write a brief but clear explanation of it.

**Programming Exercises**

1. Using the structures `parent(X, Y), male(X), and female(X), write a structure that defines mother(X, Y).`

2. Using the structures `parent(X, Y), male(X), and female(X), write a structure that defines sister(X, Y).`

3. Write a Prolog program to check if a number is prime.

4. Write a Prolog program that succeeds if the intersection of two given list parameters is empty.

5. Write a Prolog program to find the greatest common divisor of two numbers.

6. Write a Prolog program to print the Fibonacci series up to the nth term.

7. Write a Prolog program that implements an insertion sort.
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