

# **Data Structures**

# and Algorithms

ALFRED V. AHO

Bell Laboratories Murray Hill, New Jersey

#### JOHN E. HOPCROFT

Cornell University Ithaca, New York

## JEFFREY D. ULLMAN

Stanford University Stanford, California

ADDISON-WESLEY PUBLISHING COMPANY Reading, Massachusetts 

Menio Park, California London 

Amsterdam 

Don Mills, Ontario 

Sydney This book is in the ADDISON-WESLEY SERIES IN COMPUTER SCIENCE AND INFORMATION PROCESSING

Michael A. Harrison Consulting Editor

# Library of Congress Cataloging in Publication Data

Aho, Alfred V. Data structures and algorithms.

1. Data structures (Computer science) 2. Algorithms. I. Hopcroft, John E., 1939- 11. Uilman, Jeffrey D., 1942- 111. Title. QA76.9.D35A38 1982 001.64 82-11596 ISBN 0-201-00023-7

Reproduced by Addison-Wesley from camera-ready copy supplied by the authors.

## Reprinted with corrections April, 1987

Copyright © 1983 by Bell Telephone Laboratories, Incorporated.

All rights reserved. No part of this publication may be reproduced, stored in a retrieval system, or transmitted, in any form or by any means, electronic, mechanical, photocopying, recording, or otherwise, without the prior written permission of the publisher. Printed in the United States of America. Published simultaneously in Canada.

ISBN: 0-201-00023-7

# Contents

# Chapter 1 Design and Analysis of Algorithms

1.1	From Problems to Programs	ŀ
1.1	At the Date Trippe	
1.2	Abstract Data Types	13
1.3	Data Types, Data Structures, and Abstract Data Types	
14	The Running Time of a Program	16
1.5	Calculating the Running Time of a Program	
1.6	Good Programming Practice	
1.0	Super Based	
1./	Super rascal	

## Chapter 2 Basic Data Types

	••	
2.1	The Data Type "List"	
2.2	Implementation of Lists	40
2.3	Stacks	
2.4	Queues	
2.5	Mappings	61
2.6	Stacks and Recursive Procedures	

#### Chapter 3 Trees

		.75
3.1	Basic Terminology	
3.2	The ADT TREE	82
3.3	Implementations of Trees	
3.4	Binary Trees	

## Chapter 4 Basic Operations on Sets

107
109
112
115
115
117
119
122
129
135
120
130
145



#### CONTENTS

Chapter 5	Advanced Set Representation Methods
5.1 5.2 5.3 5.4 5.5 5.6	Binary Search Trees       155         Time Analysis of Binary Search Tree Operations       160         Tries       163         Balanced Tree Implementations of Sets       169         Sets with the MERGE and FIND Operations       180         An ADT with MERGE and SPLIT       189
Chapter 6	Directed Graphs
6.1 6.2 6.3 6.4 6.5 6.6 6.7	Basic Definitions198Representations for Directed Graphs199The Single-Source Shortest Paths Problem203The All-Pairs Shortest Path Problem208Traversals of Directed Graphs215Directed Acyclic Graphs219Strong Components222
Chapter 7	Undirected Graphs
7.1 7.2 7.3 7.4 7.5	Definitions       230         Minimum-Cost Spanning Trees       233         Traversals       239         Articulation Points and Biconnected Components       244         Graph Matching       246
Chapter 8	Sorting
8.1 8.2 8.3 8.4 8.5 8.6 8.7	The Internal Sorting Model
Chapter 9	Algorithm Analysis Techniques
9.1 9.2 9.3 9.4	Efficiency of Algorithms
Chapter 10	Algorithm Design Techniques
10.1 10.2 10.3 10.4 10.5	Divide-and-Conquer Algorithms306Dynamic Programming311Greedy Algorithms321Backtracking324Local Search Algorithms336

х

CONTENTS
----------

Chapter 11	Data Structures and Algorithms for External Storage	
Unapter 11	Dura Street.	347
11.1	A Model of External Computation	349
11.2	External Sorting	361
11.3	Storing Information in Files	368
11.4	External Search Trees	
Chapter 12	Memory Management	
Chapter 14	The second secon	378
12.1	The Issues in Memory Management	382
12.2	Managing Equal-Sized Blocks	384
12.3	Garbage Collection Algorithms for Equal-Sized Blocks	392
12.4	Storage Allocation for Objects with Mixed Sizes	400
12.5	Buddy Systems	.404
12.6	Storage Compaction	
	Bibliography	411
	Index	419

xi

## CHAPTER 3

# Trees

A tree imposes a hierarchical structure on a collection of items. Familiar examples of trees are genealogies and organization charts. Trees are used to help analyze electrical circuits and to represent the structure of mathematical formulas. Trees also arise naturally in many different areas of computer science. For example, trees are used to organize information in database systems and to represent the syntactic structure of source programs in compilers. Chapter 5 describes applications of trees in the representation of data. Throughout this book, we shall use many different variants of trees. In this chapter we introduce the basic definitions and present some of the more common tree operations. We then describe some of the more frequently used data structures for trees that can be used to support these operations efficiently.

#### 3.1 Basic Terminology

A tree is a collection of elements called *nodes*, one of which is distinguished as a *root*, along with a relation ("parenthood") that places a hierarchical structure on the nodes. A node, like an element of a list, can be of whatever type we wish. We often depict a node as a letter, a string, or a number with a circle around it. Formally, a *tree* can be defined recursively in the following manner.

- 1. A single node by itself is a tree. This node is also the root of the tree.
- 2. Suppose *n* is a node and  $T_1, T_2, \ldots, T_k$  are trees with roots  $n_1, n_2, \ldots, n_k$ , respectively. We can construct a new tree by making *n* be the parent of nodes  $n_1, n_2, \ldots, n_k$ . In this tree *n* is the root and  $T_1, T_2, \ldots, T_k$  are the subtrees of the root. Nodes  $n_1, n_2, \ldots, n_k$  are called the *children* of node *n*.

Sometimes, it is convenient to include among trees the *null tree*, a "tree" with no nodes, which we shall represent by  $\Lambda$ .

**Example 3.1.** Consider the table of contents of a book, as suggested by Fig. 3.1(a). This table of contents is a tree. We can redraw it in the manner shown in Fig. 3.1(b). The parent-child relationship is depicted by a line. Trees are normally drawn top-down as in Fig. 3.1(b), with the parent above the child.

The root, the node called "Book," has three subtrees with roots corresponding to the chapters C1, C2, and C3. This relationship is represented by the lines downward from Book to C1, C2, and C3. Book is the parent of C1, C2, and C3, and these three nodes are the children of Book.



Fig. 3.1. A table of contents and its tree representation.

The third subtree, with root C3, is a tree of a single node, while the other two subtrees have a nontrivial structure. For example, the subtree with root C2 has three subtrees, corresponding to the sections s2.1, s2.2, and s2.3; the last two are one-node trees, while the first has two subtrees corresponding to the subsections s2.1.1 and s2.1.2.  $\Box$ 

Example 3.1 is typical of one kind of data that is best represented as a tree. In this example, the parenthood relationship stands for containment; a parent node is comprised of its children, as Book is comprised of C1, C2, and C3. Throughout this book we shall encounter a variety of other relationships that can be represented by parenthood in trees.

If  $n_1, n_2, \ldots, n_k$  is a sequence of nodes in a tree such that  $n_i$  is the parent of  $n_{i+1}$  for  $1 \le i < k$ , then this sequence is called a *path* from node  $n_1$  to node  $n_k$ . The *length* of a path is one less than the number of nodes in the path. Thus there is a path of length zero from every node to itself. For example, in Fig. 3.1 there is a path of length two, namely (C2, s2.1, s2.1.2) from C2 to s2.1.2.

If there is a path from node a to node b, then a is an *ancestor* of b, and b is a *descendant* of a. For example, in Fig. 3.1, the ancestors of s2.1, are itself, C2, and Book, while its descendants are itself, s2.1.1, and s2.1.2. Notice that any node is both an ancestor and a descendant of itself.

An ancestor or descendant of a node, other than the node itself, is called a *proper* ancestor or *proper* descendant, respectively. In a tree, the root is the only node with no proper ancestors. A node with no proper descendants is called a *leaf*. A subtree of a tree is a node, together with all its descendants.

The *height* of a node in a tree is the length of a longest path from the node to a leaf. In Fig. 3.1 node C1 has height 1, node C2 height 2, and node C3 height 0. The *height of a tree* is the height of the root. The *depth* of a node is the length of the unique path from the root to that node.

#### 3.1 BASIC TERMINOLOGY

#### The Order of Nodes

The children of a node are usually ordered from left-to-right. Thus the two trees of Fig. 3.2 are different because the two children of node a appear in a different order in the two trees. If we wish explicitly to ignore the order of children, we shall refer to a tree as an *unordered* tree.



Fig. 3.2. Two distinct (ordered) trees.

The "left-to-right" ordering of *siblings* (children of the same node) can be extended to compare any two nodes that are not related by the ancestor-descendant relationship. The relevant rule is that if a and b are siblings, and a is to the left of b, then all the descendants of a are to the left of all the descendants of b.

**Example 3.2.** Consider the tree in Fig. 3.3. Node 8 is to the right of node 2, to the left of nodes 9, 6, 10, 4, and 7, and neither left nor right of its ancestors 1, 3, and 5.



Fig. 3.3. A tree.

A simple rule, given a node n, for finding those nodes to its left and those to its right, is to draw the path from the root to n. All nodes branching off to the left of this path, and all descendants of such nodes, are to the left of n. All nodes and descendants of nodes branching off to the right are to the right of n.  $\Box$ 

#### Preorder, Postorder, and Inorder

78

There are several useful ways in which we can systematically order all nodes of a tree. The three most important orderings are called preorder, inorder and postorder; these orderings are defined recursively as follows.

- If a tree T is null, then the empty list is the preorder, inorder and postorder listing of T.
- If T consists a single node, then that node by itself is the preorder, inorder, and postorder listing of T.

Otherwise, let T be a tree with root n and subtrees  $T_1, T_2, \ldots, T_k$ , as suggested in Fig. 3.4.



Fig. 3.4. Tree T.

- 1. The preorder listing (or preorder traversal) of the nodes of T is the root n of T followed by the nodes of  $T_1$  in preorder, then the nodes of  $T_2$  in preorder, and so on, up to the nodes of  $T_k$  in preorder.
- 2. The *inorder listing* of the nodes of T is the nodes of  $T_1$  in inorder, followed by node n, followed by the nodes of  $T_2, \ldots, T_k$ , each group of nodes in inorder.
- 3. The postorder listing of the nodes of T is the nodes of  $T_1$  in postorder, then the nodes of  $T_2$  in postorder, and so on, up to  $T_k$ , all followed by node n.

Figure 3.5(a) shows a sketch of a procedure to list the nodes of a tree in preorder. To make it a postorder procedure, we simply reverse the order of steps (1) and (2). Figure 3.5(b) is a sketch of an inorder procedure. In each case, we produce the desired ordering of the tree by calling the appropriate procedure on the root of the tree.

**Example 3.3.** Let us list the tree of Fig. 3.3 in preorder. We first list 1 and then call PREORDER on the first subtree of 1, the subtree with root 2. This subtree is a single node, so we simply list it. Then we proceed to the second subtree of 1, the tree rooted at 3. We list 3, and then call PREORDER on the first subtree of 3. That call results in listing 5, 8, and 9, in that order.

TREES

3.1 BASIC TERMINOLOGY

procedure PREORDER ( n: node );

begin

list n;

(1) (2)

(a) PREORDER procedure.

end; { INORDER }

(b) INORDER procedure.

Fig. 3.5. Recursive ordering procedures.

Continuing in this manner, we obtain the complete preorder traversal of Fig. 3.3: 1, 2, 3, 5, 8, 9, 6, 10, 4, 7.

Similarly, by simulating Fig. 3.5(a) with the steps reversed, we can discover that the postorder of Fig. 3.3 is 2, 8, 9, 5, 10, 6, 3, 7, 4, 1. By simulating Fig. 3.5(b), we find that the inorder listing of Fig. 3.3 is 2, 1, 8, 5, 9, 3, 10, 6, 7, 4.  $\Box$ 

A useful trick for producing the three node orderings is the following. Imagine we walk around the outside of the tree, starting at the root, moving counterclockwise, and staying as close to the tree as possible; the path we have in mind for Fig. 3.3 is shown in Fig. 3.6.

For preorder, we list a node the first time we pass it. For postorder, we list a node the last time we pass it, as we move up to its parent. For inorder, we list a leaf the first time we pass it, but list an interior node the second time we pass it. For example, node 1 in Fig. 3.6 is passed the first time at the beginning, and the second time while passing through the "bay" between nodes 2 and 3. Note that the order of the leaves in the three orderings is





always the same left-to-right ordering of the leaves. It is only the ordering of the interior nodes and their relationship to the leaves that vary among the three.  $\Box$ 

#### Labeled Trees and Expression Trees

80

Often it is useful to associate a *label*, or value, with each node of a tree, in the same spirit with which we associated a value with a list element in the previous chapter. That is, the label of a node is not the name of the node, but a value that is "stored" at the node. In some applications we shall even change the label of a node, while the name of a node remains the same. A useful analogy is tree: list = label: element = node: position.

**Example 3.4.** Figure 3.7 shows a labeled tree representing the arithmetic expression (a+b) \* (a+c), where  $n_1, \ldots, n_7$  are the names of the nodes, and the labels, by convention, are shown next to the nodes. The rules whereby a labeled tree represents an expression are as follows:

- 1. Every leaf is labeled by an operand and consists of that operand alone. For example, node  $n_4$  represents the expression a.
- 2. Every interior node *n* is labeled by an operator. Suppose *n* is labeled by a binary operator  $\theta$ , such as + or \*, and that the left child represents expression  $E_1$  and the right child  $E_2$ . Then *n* represents expression  $(E_1) \theta (E_2)$ . We may remove the parentheses if they are not necessary.

For example, node  $n_2$  has operator +, and its left and right children represent the expressions a and b, respectively. Therefore,  $n_2$  represents (a)+(b), or just a+b. Node  $n_1$  represents (a+b)\*(a+c), since \* is the label

TREES

3.1 BASIC TERMINOLOGY

at  $n_1$ , and a+b and a+c are the expressions represented by  $n_2$  and  $n_3$ , respectively.  $\Box$ 





Often, when we produce the preorder, inorder, or postorder listing of a tree, we prefer to list not the node names, but rather the labels. In the case of an expression tree, the preorder listing of the labels gives us what is known as the *prefix* form of an expression, where the operator precedes its left operand and its right operand. To be precise, the prefix expression for a single operand a is a itself. The prefix expression for  $(E_1) \ \theta(E_2)$ , with  $\theta$  a binary operator, is  $\theta P_1 P_2$ , where  $P_1$  and  $P_2$  are the prefix expressions for  $E_1$  and  $E_2$ . Note that no parentheses are necessary in the prefix expression, since we can scan the prefix expression  $\theta P_1 P_2$  and uniquely identify  $P_1$  as the shortest (and only) prefix of  $P_1 P_2$  that is a legal prefix expression.

For example, the preorder listing of the labels of Fig. 3.7 is \*+ab+ac. The prefix expression for  $n_2$ , which is +ab, is the shortest legal prefix of +ab+ac.

Similarly, a postorder listing of the labels of an expression tree gives us what is known as the *postfix* (or *Polish*) representation of an expression. The expression  $(E_1) \theta (E_2)$  is represented by the postfix expression  $P_1P_2\theta$ , where  $P_1$  and  $P_2$  are the postfix representations of  $E_1$  and  $E_2$ , respectively. Again, no parentheses are necessary in the postfix representation, as we can deduce what  $P_2$  is by looking for the shortest suffix of  $P_1P_2$  that is a legal postfix expression. For example, the postfix expression for Fig. 3.7 is ab+ac+\*. If we write this expression as  $P_1P_2*$ , then  $P_2$  is ac+, the shortest suffix of ab+ac+ that is a legal postfix expression.

#### TREES

The inorder traversal of an expression tree gives the infix expression itself, but with no parentheses. For example, the inorder listing of the labels of Fig. 3.7 is a+b \* a+c. The reader is invited to provide an algorithm for traversing an expression tree and producing an infix expression with all needed pairs of parentheses.

#### **Computing Ancestral Information**

The preorder and postorder traversals of a tree are useful in obtaining ancestral information. Suppose *postorder*(n) is the position of node n in a postorder listing of the nodes of a tree. Suppose desc(n) is the number of proper descendants of node n. For example, in the tree of Fig. 3.7 the postorder numbers of nodes  $n_2$ ,  $n_4$ , and  $n_5$  are 3, 1, and 2, respectively.

The postorder numbers assigned to the nodes have the useful property that the nodes in the subtree with root n are numbered consecutively from postorder(n) - desc(n) to postorder(n). To test if a vertex x is a descendant of vertex y, all we need do is determine whether

#### $postorder(y) - desc(y) \le postorder(x) \le postorder(y)$ .

A similar property holds for preorder.

#### 3.2 The ADT TREE

In Chapter 2, lists, stacks, queues, and mappings were treated as abstract data types (ADT's). In this chapter trees will be treated both as ADT's and as data structures. One of our most important uses of trees occurs in the design of implementations for the various ADT's we study. For example, in Section 5.1, we shall see how a "binary search tree" can be used to implement abstract data types based on the mathematical model of a set, together with operations such as INSERT, DELETE, and MEMBER (to test whether an element is in a set). The next two chapters present a number of other tree implementations of various ADT's.

In this section, we shall present several useful operations on trees and show how tree algorithms can be designed in terms of these operations. As with lists, there are a great variety of operations that can be performed on trees. Here, we shall consider the following operations:

- 1. PARENT(n, T). This function returns the parent of node n in tree T. If n is the root, which has no parent,  $\Lambda$  is returned. In this context,  $\Lambda$  is a "null node," which is used as a signal that we have navigated off the tree.
- 2. LEFTMOST\_CHILD(n, T) returns the leftmost child of node n in tree T, and it returns  $\Lambda$  if n is a leaf, which therefore has no children.
- 3. RIGHT\_SIBLING(n, T) returns the right sibling of node n in tree T, defined to be that node m with the same parent p as n such that m lies immediately to the right of n in the ordering of the children of p. For example, for the tree in Fig. 3.7, LEFTMOST\_CHILD $(n_2) = n_4$ ; RIGHT\_SIBLING $(n_4) = n_5$ , and RIGHT\_SIBLING  $(n_5) = \Lambda$ .

3.2 THE ADT TREE

- 4. LABEL(n, T) returns the label of node n in tree T. We do not, however, require labels to be defined for every tree.
- 5. CREATE $i(v, T_1, T_2, \ldots, T_i)$  is one of an infinite family of functions, one for each value of  $i = 0, 1, 2, \ldots$  CREATEi makes a new node r with label v and gives it i children, which are the roots of trees  $T_1, T_2, \ldots, T_i$ , in order from the left. The tree with root r is returned. Note that if i = 0, then r is both a leaf and the root.
- 6. ROOT(T) returns the node that is the root of tree T, or  $\Lambda$  if T is the null tree.
- 7. MAKENULL(T) makes T be the null tree.

**Example 3.5.** Let us write both recursive and nonrecursive procedures to take a tree and list the labels of its nodes in preorder. We assume that there are data types node and TREE already defined for us, and that the data type TREE is for trees with labels of the type labeltype. Figure 3.8 shows a recursive procedure that, given node n, lists the labels of the subtree rooted at n in preorder. We call PREORDER(ROOT(T)) to get a preorder listing of tree T.

Fig. 3.8. A recursive preorder listing procedure.

We shall also develop a nonrecursive procedure to print a tree in preorder. To find our way around the tree, we shall use a stack S, whose type STACK is really "stack of nodes." The basic idea underlying our algorithm is that when we are at a node n, the stack will hold the path from the root to n, with the root at the bottom of the stack and node n at the top.<sup>†</sup>

<sup>&</sup>lt;sup>†</sup> Recall our discussion of recursion in Section 2.6 in which we illustrated how the implementation of a recursive procedure involves a stack of activation records. If we examine Fig. 3.8, we can observe that when PREORDER(n) is called, the active procedure calls, and therefore the stack of activation records, correspond to the calls of PREORDER for all the ancestors of n. Thus our nonrecursive procedure procedure, like the example in Section 2.6, models closely the way the recursive procedure is implemented.

One way to perform a nonrecursive preorder traversal of a tree is given by the program NPREORDER shown in Fig. 3.9. This program has two modes of operation. In the first mode it descends down the leftmost unexplored path in the tree, printing and stacking the nodes along the path, until it reaches a leaf.

The program then enters the second mode of operation in which it retreats back up the stacked path, popping the nodes of the path off the stack, until it encounters a node on the path with a right sibling. The program then reverts back to the first mode of operation, starting the descent from that unexplored right sibling.

The program begins in mode one at the root and terminates when the stack becomes empty. The complete program is shown in Fig. 3.9.

#### **3.3 Implementations of Trees**

In this section we shall present several basic implementations for trees and discuss their capabilities for supporting the various tree operations introduced in Section 3.2.

#### An Array Representation of Trees

Let T be a tree in which the nodes are named  $1, 2, \ldots, n$ . Perhaps the simplest representation of T that supports the PARENT operation is a linear array A in which entry A[i] is a pointer or a cursor to the parent of node i. The root of T can be distinguished by giving it a null pointer or a pointer to itself as parent. In Pascal, pointers to array elements are not feasible, so we shall have to use a cursor scheme where A[i] = j if node j is the parent of node i, and A[i] = 0 if node i is the root.

This representation uses the property of trees that each node has a unique parent. With this representation the parent of a node can be found in constant time. A path going up the tree, that is, from node to parent to parent, and so on, can be traversed in time proportional to the number of nodes on the path. We can also support the LABEL operator by adding another array L, such that L[i] is the label of node i, or by making the elements of array A be records consisting of an integer (cursor) and a label.

**Example 3.6.** The tree of Fig. 3.10(a) has the parent representation given by the array A shown in Fig. 3.10(b).  $\Box$ 

The parent pointer representation does not facilitate operations that require child-of information. Given a node n, it is expensive to determine the children of n, or the height of n. In addition, the parent pointer representation does not specify the order of the children of a node. Thus, operations like LEFTMOST\_CHILD and RIGHT\_SIBLING are not well defined. We could impose an artificial order, for example, by numbering the children of each node after numbering the parent, and numbering the children in

```
3.3 IMPLEMENTATIONS OF TREES
        procedure NPREORDER ( T: TREE );
           { nonrecursive preorder traversal of tree T }
           var
               m: node; { a temporary }
               S: STACK; { stack of nodes holding path from the root
                   to the parent TOP(S) of the "current" node m }
           begin
               { initialize }
               MAKENULL(S);
               m := \operatorname{ROOT}(T);
               while true do
                   if m <> \Lambda then begin
                       print(LABEL(m, T));
                       PUSH(m, S);
                       \{ explore leftmost child of m \}
                       m := LEFTMOST_CHILD(m, T)
                   end
                   else begin
                       { exploration of path on stack
                           is now complete }
                       if EMPTY(S) then
                           return:
                       { explore right sibling of node
                          on top of stack }
                       m := RIGHT_SIBLING(TOP(S), T);
                       POP(S)
                   end
           end; { NPREORDER }
```

Fig. 3.9. A nonrecursive preorder procedure.

increasing order from left to right. On that assumption, we have written the function RIGHT\_SIBLING in Fig. 3.11, for types node and TREE that are defined as follows:

type
 node = integer;
 TREE = array [1..maxnodes] of node;

For this implementation we assume the null node  $\Lambda$  is represented by 0.



#### 3.3 IMPLEMENTATIONS OF TREES

#### **Representation of Trees by Lists of Children**

An important and useful way of representing trees is to form for each node a list of its children. The lists can be represented by any of the methods suggested in Chapter 2, but because the number of children each node may have can be variable, the linked-list representations are often more appropriate.

Figure 3.12 suggests how the tree of Fig. 3.10(a) might be represented. There is an array of header cells, indexed by nodes, which we assume to be numbered 1, 2, ..., 10. Each header points to a linked list of "elements," which are nodes. The elements on the list headed by *header*[i] are the children of node i; for example, 9 and 10 are the children of 3.



Fig. 3.12. A linked-list representation of a tree.

Let us first develop the data structures we need in terms of an abstract data type LIST (of nodes), and then give a particular implementation of lists and see how the abstractions fit together. Later, we shall see some of the simplifications we can make. We begin with the following type declarations:

#### type

```
node = integer;
LIST = { appropriate definition for list of nodes };
position = { appropriate definition for positions in lists };
TREE = record
    header: array [1..maxnodes] of LIST;
    labels: array [1..maxnodes] of labeltype;
    root: node
end;
```

We assume that the root of each tree is stored explicitly in the *root* field. Also, 0 is used to represent the null node.

Figure 3.13 shows the code for the LEFTMOST\_CHILD operation. The reader should write the code for the other operations as exercises.

function LEFTMOST\_CHILD ( n: node; T: TREE ) : node;
{ returns the leftmost child of node n of tree T }
var
L: LIST; { shorthand for the list of n's children }
begin
L := T.header[n];
if EMPTY(L) then { n is a leaf }
return (0)
else

return (RETRIEVE(FIRST(L), L))
end; { LEFTMOST\_CHILD }

#### Fig. 3.13. Function to find leftmost child.

Now let us choose a particular implementation of lists, in which both LIST and position are integers, used as cursors into an array *cellspace* of records:

var

cellspace: array [1..maxnodes] of record node: integer; next: integer end;

To simplify, we shall not insist that lists of children have header cells. Rather, we shall let *T.header* [n] point directly to the first cell of the list, as is suggested by Fig. 3.12. Figure 3.14(a) shows the function LEFTMOST\_CHILD of Fig. 3.13 rewritten for this specific implementation. Figure 3.14(b) shows the operator PARENT, which is more difficult to write using this representation of lists, since a search of all lists is required to determine on which list a given node appears.

#### The Leftmost-Child, Right-Sibling Representation

The data structure described above has, among other shortcomings, the inability to create large trees from smaller ones, using the CREATE*i* operators. The reason is that, while all trees share *cellspace* for linked lists of children, each has its own array of headers for its nodes. For example, to implement CREATE2( $\nu$ ,  $T_1$ ,  $T_2$ ) we would have to copy  $T_1$  and  $T_2$  into a third tree and add a new node with label  $\nu$  and two children — the roots of  $T_1$  and  $T_2$ .

If we wish to build trees from smaller ones, it is best that the representation of nodes from all trees share one area. The logical extension of Fig. 3.12 is to replace the header array by an array *nodespace* consisting of records with

```
function LEFTMOST_CHILD ( n: node; T: TREE ) : node;
    { returns the leftmost child of node n on tree T }
    var
        L: integer; { a cursor to the beginning of the list of n's children }
    begin
        L := T.header[n];
        if L = 0 then \{n \text{ is a leaf}\}
            return (0)
        else
            return (cellspace[L].node)
    end; { LEFTMOST_CHILD }
          (a) The function LEFTMOST_CHILD.
function PARENT ( n: node; T: TREE ) : node;
    { returns the parent of node n in tree T }
    var
        p: node; { runs through possible parents of n }
        i: position; { runs down list of p's children }
    begin
        for p := 1 to maxnodes do begin
            i := T.header[p];
            while i \le 0 do { see if n is among children of p }
                if cellspace[i].node = n then
                    return (p)
                else
                    i := cellspace[i].next
        end;
        return (0) { return null node if parent not found }
    end; { PARENT }
                (b) The function PARENT.
```

Fig. 3.14. Two functions using linked-list representation of trees.

two fields *label* and *header*. This array will hold headers for all nodes of all trees. Thus, we declare

```
var
nodespace: array [1..maxnodes] of record
label: labeltype;
header: integer { cursor to cellspace }
end;
```

Then, since nodes are no longer named 1, 2,  $\ldots$ , *n*, but are represented by

arbitrary indices in *nodespace*, it is no longer feasible for the field *node* of *cellspace* to represent the "number" of a node; rather, *node* is now a cursor into *nodespace*, indicating the position of that node. The type TREE is simply a cursor into *nodespace*, indicating the position of the root.

**Example 3.7.** Figure 3.15(a) shows a tree, and Fig. 3.15(b) shows the data structure where we have placed the nodes labeled A, B, C, and D arbitrarily in positions 10, 5, 11, and 2 of *nodespace*. We have also made arbitrary choices for the cells of *cellspace* used for lists of children.  $\Box$ 



Fig. 3.15. Another linked-list structure for trees.

The structure of Fig. 3.15(b) is adequate to merge trees by the CREATEi operations. This data structure can be significantly simplified, however. First, observe that the chains of *next* pointers in *cellspace* are really right-sibling pointers.

Using these pointers, we can obtain leftmost children as follows. Suppose cellspace[i].node = n. (Recall that the "name" of a node, as opposed to its label, is in effect its index in nodespace, which is what cellspace[i].node gives us.) Then nodespace[n].header indicates the cell for the leftmost child of n in cellspace, in the sense that the node field of that cell is the name of that node in nodespace.

We can simplify matters if we identify a node not with its index in

#### 3.3 IMPLEMENTATIONS OF TREES

*nodespace*, but with the index of the cell in *cellspace* that represents it as a child. Then, the *next* pointers of *cellspace* truly point to right siblings, and the information contained in the *nodespace* array can be held by introducing a field *leftmost\_child* in *cellspace*. The datatype TREE becomes an integer used as a cursor to *cellspace* indicating the root of the tree. We declare *cellspace* to have the following structure.

var
 cellspace: array [1..maxnodes] of record
 label: labeltype;
 leftmost\_child: integer;
 right\_sibling: integer
end;

**Example 3.8.** The tree of Fig. 3.15(a) is represented in our new data structure in Fig. 3.16. The same arbitrary indices as in Fig. 3.15(b) have been used for the nodes.  $\Box$ 



Fig. 3.16. Leftmost-child, right-sibling representation of a tree.

All operations but PARENT are straightforward to implement in the leftmost-child, right-sibling representation. PARENT requires searching the entire *cellspace*. If we need to perform the PARENT operation efficiently, we can add a fourth field to *cellspace* to indicate the parent of a node directly.

As an example of a tree operation written to use the leftmost-child, right-

TREES

sibling structure as in Fig. 3.16, we give the function CREATE2 in Fig. 3.17. We assume that unused cells are linked in an available space list, headed by *avail*, and that available cells are linked by their right-sibling fields. Figure 3.18 shows the old (solid) and the new (dashed) pointers.

```
return (temp)
end; { CREATE2 }
```

92





# Fig. 3.18. Pointer changes produced by CREATE2.

Alternatively, we can use less space but more time if we put in the right-

#### 3.3 IMPLEMENTATIONS OF TREES

sibling field of the rightmost child a pointer to the parent, in place of the null pointer that would otherwise be there. To avoid confusion, we need a bit in every cell indicating whether the right-sibling field holds a pointer to the right sibling or to the parent.

Given a node, we find its parent by following right-sibling pointers until we find one that is a parent pointer. Since all siblings have the same parent, we thereby find our way to the parent of the node we started from. The time required to find a node's parent in this representation depends on the number of siblings a node has.

#### 3.4 Binary Trees

The tree we defined in Section 3.1 is sometimes called an *ordered*, *oriented* tree because the children of each node are ordered from left-to-right, and because there is an oriented path (path in a particular direction) from every node to its descendants. Another useful, and quite different, notion of "tree" is the *binary tree*, which is either an empty tree, or a tree in which every node has either no children, a *left child*, a *right child*, or both a left and a right child. The fact that each child in a binary tree is designated as a left child or as a right child makes a binary tree different from the ordered, oriented tree of Section 3.1.

**Example 3.9.** If we adopt the convention that left children are drawn extending to the left, and right children to the right, then Fig. 3.19 (a) and (b) represent two different binary trees, even though both "look like" the ordinary (ordered, oriented) tree of Fig. 3.20. However, let us emphasize that Fig. 3.19(a) and (b) are not the same binary tree, nor are either in any sense equal to Fig. 3.20, for the simple reason that binary trees are not directly comparable with ordinary trees. For example, in Fig. 3.19(a), 2 is the left child of 1, and 1 has no right child, while in Fig. 3.19(b), 1 has no left child but has 2 as a right child. In either binary tree, 3 is the left child of 2, and 4 is 2's right child.  $\square$ 

The preorder and postorder listings of a binary tree are similar to those of an ordinary tree given on p. 78. The inorder listing of the nodes of a binary tree with root n, left subtree  $T_1$  and right subtree  $T_2$  is the inorder listing of  $T_1$ followed by n followed by the inorder listing of  $T_2$ . For example, 35241 is the inorder listing of the nodes of Fig. 3.19(a).

#### **Representing Binary Trees**

A convenient data structure for representing a binary tree is to name the nodes  $1, 2, \ldots, n$ , and to use an array of records declared

- var
  - cellspace: array [1...maxnodes] of record leftchild: integer; rightchild: integer end;



Fig. 3.19. Two binary trees.



Fig. 3.20. An "ordinary" tree.

The intention is that cellspace[i].leftchild is the left child of node *i*, and *rightchild* is analogous. A value of 0 in either field indicates the absence of a child.

**Example 3.10.** The binary tree of Fig. 3.19(a) can be represented as shown in Fig. 3.21.  $\square$ 

#### An Example: Huffman Codes

94

Let us give an example of how binary trees can be used as a data structure. The particular problem we shall consider is the construction of "Huffman codes." Suppose we have messages consisting of sequences of characters. In each message, the characters are independent and appear with a known

	leftchild	rightchild
1	2	0
2	3	4
3	0	5
4	0	0
5	0	0

Fig. 3.21. Representation of a binary tree.

probability in any given position; the probabilities are the same for all positions. As an example, suppose we have a message made from the five characters a, b, c, d, e, which appear with probabilities .12, .4, .15, .08, .25, respectively.

We wish to encode each character into a sequence of 0's and 1's so that no code for a character is the prefix of the code for any other character. This *prefix property* allows us to decode a string of 0's and 1's by repeatedly deleting prefixes of the string that are codes for characters.

**Example 3.11.** Figure 3.22 shows two possible codes for our five symbol alphabet. Clearly Code 1 has the prefix property, since no sequence of three bits can be the prefix of another sequence of three bits. The decoding algorithm for Code 1 is simple. Just "grab" three bits at a time and translate each group of three into a character. Of course, sequences 101, 110, and 111 are impossible, if the string of bits really codes characters according to Code 1. For example, if we receive 001010011 we know the original message was *bcd*.

Symbol	Probability	Code 1	Code 2
a	.12	000	000
b	.40	001	11
с	.15	010	01
d	.08	011	001
е	.25	100	10

#### Fig. 3.22. Two binary codes.

It is easy to check that Code 2 also has the prefix property. We can decode a string of bits by repeatedly "grabbing" prefixes that are codes for characters and removing them, just as we did for Code 1. The only difference is that here, we cannot slice up the entire sequence of bits at once, because whether we take two or three bits for a character depends on the bits. For example, if a string begins 1101001, we can again be sure that the characters coded were *bcd*. The first two bits, 11, must have come from *b*, so we can

remove them and worry about 01001. We then deduce that the bits 01 came from c, and so on.  $\Box$ 

The problem we face is: given a set of characters and their probabilities, find a code with the prefix property such that the average length of a code for a character is a minimum. The reason we want to minimize the average code length is to compress the length of an average message. The shorter the average code for a character is, the shorter the length of the encoded message. For example, Code I has an average code length of 3. This is obtained by multiplying the length of the code for each symbol by the probability of occurrence of that symbol. Code 2 has an average length of 2.2, since symbols a and d, which together appear 20% of the time, have codes of length three, and the other symbols have codes of length two.

Can we do better than Code 2? A complete answer to this question is to exhibit a code with the prefix property having an average length of 2.15. This is the best possible code for these probabilities of symbol occurrences. One technique for finding optimal prefix codes is called *Huffman's algorithm*. It works by selecting two characters a and b having the lowest probabilities and replacing them with a single (imaginary) character, say x, whose probability of occurrence is the sum of the probabilities for a and b. We then find an optimal prefix code for this smaller set of characters, using this procedure recursively. The code for the original character set is obtained by using the code for x with a 0 appended as the code for a and with a 1 appended as a code for b.

We can think of prefix codes as paths in binary trees. Think of following a path from a node to its left child as appending a 0 to a code, and proceeding from a node to its right child as appending a 1. If we label the leaves of a binary tree by the characters represented, we can represent any prefix code as a binary tree. The prefix property guarantees no character can have a code that is an interior node, and conversely, labeling the leaves of any binary tree with characters gives us a code with the prefix property for these characters.

**Example 3.12.** The binary trees for Code 1 and Code 2 of Fig. 3.22 are shown in Fig. 3.23(a) and (b), respectively.  $\Box$ 

We shall implement Huffman's algorithm using a *forest* (collection of trees), each of which has its leaves labeled by characters whose codes we desire to select and whose roots are labeled by the sum of the probabilities of all the leaf labels. We call this sum the *weight* of the tree. Initially, each character is in a one-node tree by itself, and when the algorithm ends, there will be only one tree, with all the characters at its leaves. In this tree, the path from the root to any leaf represents the code for the label of that leaf, according to the left = 0, right = 1 scheme of Fig. 3.23.

The essential step of the algorithm is to select the two trees in the forest that have the smallest weights (break ties arbitrarily). Combine these two trees into one, whose weight is the sum of the weights of the two trees. To combine the trees we create a new node, which becomes the root and has the

3.4 BINARY TREES



Fig. 3.23. Binary trees representing codes with the prefix property.

roots of the two given trees as left and right children (which is which doesn't matter). This process continues until only one tree remains. That tree represents a code that, for the probabilities given, has the minimum possible average code length.

**Example 3.13.** The sequence of steps taken for the characters and probabilities in our running example is shown in Fig. 3.24. From Fig. 3.24(e) we see the code words for a, b, c, d, and e are 1111, 0, 110, 1110, and 10. In this example, there is only one nontrivial tree, but in general, there can be many. For example, if the probabilities of b and e were .33 and .32, then after Fig. 3.24(c) we would combine b and e, rather than attaching e to the large tree as we did in Fig. 3.24(d).  $\Box$ 

Let us now describe the needed data structures. First, we shall use an array TREE of records of the type

record

*leftchild*: integer; *rightchild*: integer; *parent*: integer **end** 

to represent binary trees. Parent pointers facilitate finding paths from leaves to roots, so we can discover the code for a character. Second, we use an array *ALPHABET* of records of type

```
record
```

symbol: char; probability: real; leaf: integer { cursor into tree }

end



Fig. 3.24. Steps in the construction of a Huffman tree.

to associate, with each symbol of the alphabet being encoded, its corresponding leaf. This array also records the probability of each character. Third, we need an array *FOREST* of records that represent the trees themselves. The type of these records is

# record weight: real; root: integer { cursor into tree } end

The initial values of all these arrays, assuming the data of Fig. 3.24(a), are shown in Fig. 3.25. A sketch of the program to build the Huffman tree is shown in Fig. 3.26.

1	.12	1	1	a	.12	1		1	0	0	0
2	.40	2	2	b	.40	2		2	0	0	0
3	.15	3	3	С	.15	3		3	0	0	0
4	.08	4	4	d	.08	4		4	0	0	0
5	.25	5	5	е	.25	5	]	5	0	0	0
	weight	root		symbol	prob- ability	leaf			left- child	right- child	parent
FOREST				A	LPHABI	ET				TREE	

Fig. 3.25. Initial contents of arrays.

(1)	while there is more then one tree in the forest do begin
(2)	i := index of the tree in <i>FOREST</i> with smallest weight;
(3)	j := index of the tree in FOREST with second smallest weight;
(4)	<pre>create a new node with left child FOREST[i].root and right child FOREST[j].root;</pre>
(5)	replace tree i in FOREST by a tree whose root is the new node and whose weight is FORESTLI weight + FORESTLI weight:
(6)	delete tree j from FOREST end;

Fig. 3.26. Sketch of Huffman tree construction.

To implement line (4) of Fig. 3.26, which increases the number of cells of the *TREE* array used, and lines (5) and (6), which decrease the number of utilized cells of *FOREST*, we shall introduce cursors *lasttree* and *lastnode*, pointing to *FOREST* and *TREE*, respectively. We assume that cells 1 to *lasttree* of *FOREST* and 1 to *lastnode* of *TREE* are occupied.<sup>†</sup> We assume that arrays of Fig. 3.25 have some declared lengths, but in what follows we omit comparisons between these limits and cursor values.

<sup>&</sup>lt;sup> $\dagger$ </sup> For the data reading phase, which we omit, we also need a cursor for the array ALPHA-BET as it fills with symbols and their probabilities.

TREES

procedure lightones ( var least, second: integer ); { sets least and second to the indices in FOREST of the trees of smallest weight. We assume lasttree  $\geq 2$ . var i: integer; **begin** { initialize *least* and *second*, considering first two trees } if FOREST[1], weight  $\leq = FOREST[2]$ , weight then **begin** least := 1; second := 2 end else **begin** least := 2; second := 1 end; { Now let i run from 3 to lasttree. At each iteration least is the tree of smallest weight among the first i trees in FOREST, and second is the next smallest of these } for i := 3 to lastree do if FOREST[i].weight < FOREST[least].weight then **begin** second := least; least := i end else if FOREST[i].weight < FOREST[second].weight then second := iend; { lightones }

function create ( lefttree, righttree: integer ) : integer; { returns new node whose left and right children are

FOREST[lefttree].root and FOREST[righttree].root }

begin

lastnode := lastnode + 1; { cell for new node is TREE[lastnode] } TREE[lastnode].leftchild := FOREST[lefttree].root; TREE[lastnode].rightchild := FOREST[righttree].root; { now enter parent pointers for new node and its children } TREE[lastnode].parent := 0; TREE[FOREST[lefttree].root].parent := lastnode; TREE[FOREST[righttree].root].parent := lastnode; return (lastnode) end; { create }

Fig. 3.27. Two procedures.

Figure 3.27 shows two useful procedures. The first implements lines (2) and (3) of Fig. 3.26 to select indices of the two trees of smallest weight. The second is the command *create* $(n_1, n_2)$  that creates a new node and makes  $n_1$  and  $n_2$  its left and right children.

Now the steps of Fig. 3.26 can be described in greater detail. A procedure Huffman, which has no input or output, but works on the global structures of Fig. 3.25, is shown in Fig. 3.28.

procedure Huffman; var *i*, *j*: integer; { the two trees of least weight in FOREST } newroot: integer; begin while *lasttree* > 1 do begin lightones(i, j); newroot := create(i, j);{ Now replace tree *i* by the tree whose root is *newroot* } FOREST[i].weight := FOREST[i].weight + FOREST[j].weight; FOREST[i].root := newroot; { next, replace tree j, which is no longer needed, by lasttree, and shrink FOREST by one } FOREST[j] := FOREST[lasttree];*lasttree* := *lasttree* - 1 end end; { Huffman }

Fig. 3.28. Huffman's algorithm.

Figure 3.29 shows the data structure of Fig. 3.25 after *lasttree* has been reduced to 3, that is, when the forest looks like Fig. 3.24(c).



Fig. 3.29. Tree data structure after two iterations.

TREES

After completing execution of the algorithm, the code for each symbol can be determined as follows. Find the symbol in the symbol field of the ALPHA-BET array. Follow the leaf field of the same record to get to a record of the TREE array; this record corresponds to the leaf for that symbol. Repeatedly follow the parent pointer from the "current" record, say for node n, to the record of the TREE array for its parent p. Remember node n, so it is possible to examine the leftchild and rightchild pointers for node p and see which is n. In the former case, print 0, in the latter print 1. The sequence of bits printed is the code for the symbol, in reverse. If we wish the bits printed in the correct order, we could push each onto a stack as we go up the tree, and then repeatedly pop the stack, printing symbols as we pop them.

#### **Pointer-Based Implementations of Binary Trees**

Instead of using cursors to point to left and right children (and parents if we wish), we can use true Pascal pointers. For example, we might declare

```
type

node = record

leftchild: t node;

rightchild: t node;

parent: t node

end
```

For example, if we used this type for nodes of a binary tree, the function *create* of Fig. 3.27 could be written as in Fig. 3.30.

```
function create ( lefttree, righttree: t node ) : t node;
var
    root: t node;
begin
    new(root);
    root t.leftchild := lefttree;
    root t.rightchild := righttree;
    root t.parent := 0;
    lefttree t.parent := root;
    righttree t.parent := root;
    return (root)
end; { create }
```

Fig. 3.30. Pointer-based implementation of binary trees.

#### EXERCISES

#### Exercises

3.1 Answer the following questions about the tree of Fig. 3.31.

- a) Which nodes are leaves?
- b) Which node is the root?
- c) What is the parent of node C?
- d) Which nodes are children of C?
- e) Which nodes are ancestors of E?
- f) Which nodes are descendants of E?
- g) What are the right siblings of D and E?
- h) Which nodes are to the left and to the right of G?
- i) What is the depth of node C?
- j) What is the height of node C?



Fig. 3.31. A tree.

- 3.2 In the tree of Fig. 3.31 how many different paths of length three are there?
- **3.3** Write programs to compute the height of a tree using each of the three tree representations of Section 3.3.
- 3.4 List the nodes of Fig. 3.31 in
  - a) preorder,

- b) postorder, and
- c) inorder.
- 3.5 If m and n are two different nodes in the same tree, show that exactly one of the following statements is true:
  - a) m is to the left of n
  - b) m is to the right of n
  - c) m is a proper ancestor of n
  - d) m is a proper descendant of n.
- **3.6** Place a check in row i and column j if the two conditions represented by row i and column j can occur simultaneously.

	preorder(n) < preorder(m)	inorder(n) < inorder(m)	<pre>postorder(n) &lt; postorder(m)</pre>
n is to the left of $m$			
n is to the right of m			
<i>n</i> is a proper ancestor of <i>m</i>			
n is a proper descendant of $m$			

For example, put a check in row 3 and column 2 if you believe that n can be a proper ancestor of m and at the same time n can precede m in inorder.

- **3.7** Suppose we have arrays PREORDER[n], INORDER[n], and POSTORDER[n] that give the preorder, inorder, and postorder positions, respectively, of each node n of a tree. Describe an algorithm that tells whether node i is an ancestor of node j, for any pair of nodes i and j. Explain why your algorithm works.
- \*3.8 We can test whether a node m is a proper ancestor of a node n by testing whether m precedes n in X-order but follows n in Y-order, where X and Y are chosen from {pre, post, in}. Determine all those pairs X and Y for which this statement holds.
- 3.9 Write programs to traverse a binary tree in
  - a) preorder,
  - b) postorder,
  - c) inorder.
#### EXERCISES

- **3.10** The *level-order* listing of the nodes of a tree first lists the root, then all nodes of depth 1, then all nodes of depth 2, and so on. Nodes at the same depth are listed in left-to-right order. Write a program to list the nodes of a tree in level-order.
- 3.11 Convert the expression ((a+b) + c \* (d+e) + f) \* (g+h) to a
  - a) prefix expression
  - b) postfix expression.
- 3.12 Draw tree representations for the prefix expressions
  - a) \*a + b \* c + de
  - b) \*a + \*b + cde
- **3.13** Let T be a tree in which every nonleaf node has two children. Write a program to convert
  - a) a preorder listing of T into a postorder listing,
  - b) a postorder listing of T into a preorder listing,
  - c) a preorder listing of T into an inorder listing.
- 3.14 Write a program to evaluate
  - a) preorder
  - b) postorder

arithmetic expressions.

- **3.15** We can define a binary tree as an ADT with the binary tree structure as a mathematical model and with operations such as LEFTCHILD(n), RIGHTCHILD(n), PARENT(n), and NULL(n). The first three operations return the left child, the right child, and the parent of node n ( $\Lambda$  if there is none) and the last returns true if and only if n is  $\Lambda$ . Implement these procedures using the binary tree representation of Fig. 3.21.
- **3.16** Implement the seven tree operations of Section 3.2 using the following tree implementations:
  - a) parent pointers
  - b) lists of children
  - c) leftmost-child, right-sibling pointers.
- **3.17** The *degree* of a node is the number of children it has. Show that in any binary tree the number of leaves is one more than the number of nodes of degree two.
- **3.18** Show that the maximum number of nodes in a binary tree of height h is  $2^{h+1}-1$ . A binary tree of height h with the maximum number of nodes is called a *full* binary tree.

#### TREES

- **\*3.19** Suppose trees are implemented by leftmost-child, right-sibling and parent pointers. Give nonrecursive preorder, postorder, and inorder traversal algorithms that do not use "states" or a stack, as Fig. 3.9 does.
- **3.20** Suppose characters a, b, c, d, e, f have probabilities .07, .09, .12, .22, .23, .27, respectively. Find an optimal Huffman code and draw the Huffman tree. What is the average code length?
- \*3.21 Suppose T is a Huffman tree, and that the leaf for symbol a has greater depth than the leaf for symbol b. Prove that the probability of symbol b is no less than that of a.
- \*3.22 Prove that Huffman's algorithm works, i.e., it produces an optimal code for the given probabilities. *Hint*: Use Exercise 3.21.

#### **Bibliographic Notes**

Berge [1958] and Harary [1969] discuss the mathematical properties of trees. Knuth [1973] and Nievergelt [1974] contain additional information on binary search trees. Many of the works on graphs and applications referenced in Chapter 6 also cover material on trees.

The algorithm given in Section 3.4 for finding a tree with a minimal weighted path length is from Huffman [1952]. Parker [1980] gives some more recent explorations into that algorithm.

# **CHAPTER 5**

# Advanced Set Representation Methods

This chapter introduces data structures for sets that permit more efficient implementation of common collections of set operations than those of the previous chapter. These structures, however, are more complex and are often only appropriate for large sets. All are based on various kinds of trees, such as binary search trees, tries, and balanced trees.

#### 5.1 Binary Search Trees

We shall begin with binary search trees, a basic data structure for representing sets whose elements are ordered by some linear order. We shall, as usual, denote that order by <. This structure is useful when we have a set of elements from a universe so large that it is impractical to use the elements of the set themselves as indices into arrays. An example of such a universe would be the set of possible identifiers in a Pascal program. A binary search tree can support the set operations INSERT, DELETE, MEMBER, and MIN, taking  $O(\log n)$  steps per operation on the average for a set of n elements.

A binary search tree is a binary tree in which the nodes are labeled with elements of a set. The important property of a binary search tree is that all elements stored in the left subtree of any node x are all less than the element stored at x, and all elements stored in the right subtree of x are greater than the element stored at x. This condition, called the binary search tree property, holds for every node of a binary search tree, including the root.

Figure 5.1 shows two binary search trees representing the same set of integers. Note the interesting property that if we list the nodes of a binary search tree in inorder, then the elements stored at those nodes are listed in sorted order.

Suppose a binary search tree is used to represent a set. The binary search tree property makes testing for membership in the set simple. To determine whether x is a member of the set, first compare x with the element r at the root of the tree. If x = r we are done and the answer to the membership query is "true." If x < r, then by the binary search tree property, x can only



Fig. 5.1. Two binary search trees.

be a descendant of the left child of the root, if x is present at all.<sup>†</sup> Similarly, if x > r, then x could only be at a descendant of the right child of the root.

We shall write a simple recursive function MEMBER(x, A) to implement this membership test. We assume the elements of the set are of an unspecified type that will be called elementtype. For convenience, we assume elementtype is a type for which < and = are defined. If not, we must define functions LT(a, b) and EQ(a, b), where a and b are of type elementtype, such that LT(a, b) is true if and only if a is "less than" b, and EQ(a, b) is true if and only if a and b are the same.

The type for nodes consists of an element and two pointers to other nodes:

#### type

nodetype = record
 element: elementtype;
 leftchild, rightchild: t nodetype
end;

Then we can define the type SET as a pointer to a node, which we take to be the root of the binary search tree representing the set. That is:

<sup>†</sup> Recall the left child of the root is a descendant of itself, so we have not ruled out the possibility that x is at the left child of the root.

5.1 BINARY SEARCH TREES

SET = t nodetype;

Now we can specify fully the function MEMBER, in Fig. 5.2. Notice that since SET and "pointer to nodetype" are synonymous, MEMBER can call itself on subtrees as if those subtrees represented sets. In effect, the set can be subdivided into the subset of members less than x and the subset of members greater than x.

```
function MEMBER ( x: elementtype; A: SET ) : boolean;
{ returns true if x is in A, false otherwise }
begin
    if A = nil then
        return (false) { x is never in Ø }
    else if x = A t.element then
        return (true)
    else if x < A t.element then
        return (MEMBER(x, A t.leftchild))
    else { x > A t.element }
        return (MEMBER(x, A t.rightchild))
end; { MEMBER }
```

Fig. 5.2. Testing membership in a binary search tree.

The procedure INSERT(x, A), which adds element x to set A, is also easy to write. The first action INSERT must do is test whether A = nil, that is, whether the set is empty. If so, we create a new node to hold x and make A point to it. If the set is not empty, we search for x more or less as MEMBER does, but when we find a nil pointer during our search, we replace it by a pointer to a new node holding x. Then x will be in the right place, namely, the place where the function MEMBER will find it. The code for INSERT is shown in Fig. 5.3.

Deletion presents some problems. First, we must locate the element x to be deleted in the tree. If x is at a leaf, we can delete that leaf and be done. However, x may be at an interior node *inode*, and if we simply deleted *inode*, we would disconnect the tree.

If *inode* has only one child, as the node numbered 14 in Fig. 5.1(b), we can replace *inode* by that child, and we shall be left with the appropriate binary search tree. If *inode* has two children, as the node numbered 10 in Fig. 5.1(a), then we must find the lowest-valued element among the descendants of the right child.<sup>†</sup> For example, in the case the element 10 is deleted from Fig. 5.1(a), we must replace it by 12, the minimum-valued descendant of

<sup>†</sup> The highest-valued node among the descendants of the left child would do as well.

```
procedure INSERT ( x: elementtype; var A: SET );
           \{ add x to set A \}
           begin
               if A = nil then begin
                   new(A);
                   At .element := x;
                   At .leftchild := nil;
                   At .rightchild := nil
               end
               else if x < A^{\dagger}.element then
                    INSERT(x, At.leftchild)
                else if x > A^{\dagger}.element then
                    INSERT (x, At.rightchild)
               { if x = A_{1,element}, we do nothing; x is already in the set }
            end; { INSERT }
           Fig. 5.3. Inserting an element into a binary search tree.
the right child of 10.
    To write DELETE, it is useful to have a function DELETEMIN(A) that
removes the smallest element from a nonempty tree and returns the value of
the element removed. The code for DELETEMIN is shown in Fig. 5.4. The
code for DELETE uses DELETEMIN and is shown in Fig. 5.5.
        function DELETEMIN ( var A: SET ) : elementtype;
            { returns and removes the smallest element from set A }
        begin
            if At leftchild = nil then begin
                { A points to the smallest element }
                 DELETEMIN := At.element;
                 A := At.rightchild
                     { replace the node pointed to by A by its right child }
             end
            else { the node pointed to by A has a left child }
                 DELETEMIN := DELETEMIN(At.leftchild)
```

end; { DELETEMIN }

Fig. 5.4. Deleting the smallest element.

**Example 5.1.** Suppose we try to delete 10 from Fig. 5.1(a). Then, in the last statement of DELETE we call DELETEMIN with argument a pointer to node 14. That pointer is the *rightchild* field in the root. That call results in another call to DELETEMIN. The argument is then a pointer to node 12; this pointer is found in the *leftchild* field of node 14. We find that 12 has no

5.1 BINARY SEARCH TREES

```
procedure DELETE ( x: elementtype; var A: SET );
    { remove x from set A }
begin
    if A <> nil then
        if x < A t.element then
             DELETE(x, At.leftchild)
        else if x > A t.element then
             DELETE(x, At.rightchild)
        { if we reach here, x is at the node pointed to by A }
        else if (A^{\dagger}.leftchild = nil) and (A^{\dagger}.rightchild = nil) then
             A := nil \{ delete the leaf holding x \}
         else if A^{\dagger}. leftchild = nil then
             A := A t.rightchild
        else if A^{\dagger}.rightchild = nil then
             A := A t.leftchild
        else { both children are present }
             At.element := DELETEMIN(At.rightchild)
end; { DELETE }
```



left child, so we return element 12 and make the left child for 14 be the right child of 12, which happens to be **nil**. Then DELETE takes the value 12 returned by DELETEMIN and replaces 10 by it. The resulting tree is shown in Fig. 5.6.  $\Box$ 



Fig. 5.6. Tree of Fig. 5.1(a) after deleting 10.

# 5.2 Time Analysis of Binary Search Tree Operations

In this section we analyze the average behavior of various binary search tree operations. We show that if we insert n random elements into an initially empty binary search tree, then the average path length from the root to a leaf is  $O(\log n)$ . Testing for membership, therefore, takes  $O(\log n)$  time.

It is easy to see that if a binary tree of n nodes is complete (all nodes, except those at the lowest level have two children), then no path has more than 1+logn nodes.<sup>†</sup> Thus, each of the procedures MEMBER, INSERT, DELETE, and DELETEMIN takes  $O(\log n)$  time. To see this, observe that they all take a constant amount of time at a node, then may call themselves recursively on at most one child. Therefore, the sequence of nodes at which calls are made forms a path from the root. Since that path is  $O(\log n)$  in length, the total time spent following the path is  $O(\log n)$ .

However, when we insert n elements in a "random" order, they do not necessarily arrange themselves into a complete binary tree. For example, if we happen to insert smallest first, in sorted order, then the resulting tree is a chain of n nodes, in which each node except the lowest has a right child but no left child. In this case, it is easy to show that, as it takes O(i) steps to insert the *i*th element, and  $\sum_{i=1}^{n} i = n(n+1)/2$ , the whole process of *n* insertions

takes  $O(n^2)$  steps, or O(n) steps per operation.

We must determine whether the "average" binary search tree of n nodes is closer to the complete tree in structure than to the chain, that is, whether the average time per operation on a "random" tree takes  $O(\log n)$  steps, O(n)steps, or something in between. As we cannot know the true frequency of insertions and deletions, or whether deleted elements have some special property (e.g., do we always delete the minimum?), we can only analyze the average path length of "random" trees if we make some assumptions. The particular assumptions we make are that trees are formed by insertions only, and all orders of the n inserted elements are equally likely.

Under these fairly natural assumptions, we can calculate P(n), the average number of nodes on the path from the root to some node (not necessarily a leaf). We assume the tree was formed by inserting n random elements into an initially empty tree. Clearly P(0) = 0 and P(1) = 1. Suppose we have a list of  $n \ge 2$  elements to insert into an empty tree. The first element on the list, call it a, is equally likely to be first, second, or nth in the sorted order. Suppose that i elements on the list are less than a, so n-i-1 are greater than a. When we build the tree, a will appear at the root, the i smaller elements will be descendants of the left child of the root, and the remaining n-i-1will be descendants of the right child. This tree is sketched in Fig. 5.7.

As all orders for the *i* small elements and for the n-i-1 large elements are equally likely, we expect that the left and right subtrees of the root will

† Recall that all logarithms are to the base 2 unless otherwise noted.



Fig. 5.7. Binary search tree.

have average path lengths P(i) and P(n-i-1), respectively. Since these elements are reached through the root of the complete tree, we must add 1 to the number of nodes on every path. Thus P(n) can be calculated by averaging, for all *i* between 0 and n-1, the sum

$$\frac{i}{n}(P(i)+1) + \frac{(n-i-1)}{n}(P(n-i-1)+1) + \frac{1}{n}$$

The first term is the average path length in the left subtree, weighted by its size. The second term is the analogous quantity for the right subtree, and the 1/n term represents the contribution of the root. By averaging the above sum for all i between 1 and n, we obtain the recurrence

$$P(n) = 1 + \frac{1}{n^2} \sum_{i=0}^{n-1} \left( iP(i) + (n-i-1) P(n-i-1) \right)$$
(5.1)

The first part of the summation (5.1),  $\sum_{i=0}^{n-1} iP(i)$ , can be made identical to the second part  $\sum_{i=0}^{n-1} (n-i-1)P(n-i-1)$  if we substitute *i* for n-i-1 in the second part. Also, the term for i=0 in the summation  $\sum_{i=0}^{n-1} iP(i)$  is zero, so we

can begin the summation at 1. Thus (5.1) can be written

$$P(n) = 1 + \frac{2}{n^2} \sum_{i=1}^{n-1} i P(i) \quad \text{for } n \ge 2$$
 (5.2)

We shall show by induction on n, starting at n=1, that  $P(n) \le 1 + 4\log n$ . Surely this statement is true for n = 1, since P(1) = 1. Suppose it is true for all i < n. Then by (5.2)

$$P(n) \le 1 + \frac{2}{n^2} \sum_{i=1}^{n-1} (4i \log i + i)$$

$$\leq 1 + \frac{2}{n^2} \sum_{i=1}^{n-1} 4i \log i + \frac{2}{n^2} \sum_{i=1}^{n-1} i$$
  
$$\leq 2 + \frac{8}{n^2} \sum_{i=1}^{n-1} i \log i$$
(5.3)

The last step is justified, since  $\sum_{i=1}^{n-1} i \le n^2/2$ , and therefore, the last term of the

second line is at most 1. We shall divide the terms in the summation of (5.3) into two parts, those for  $i \leq \lfloor n/2 \rfloor - 1$ , which do not exceed  $i \log(n/2)$ , and those for  $i > \lfloor n/2 \rfloor - 1$ , which do not exceed  $i \log n$ . Thus (5.3) can be rewritten

$$P(n) \le 2 + \frac{8}{n^2} \left[ \sum_{i=1}^{\lfloor n/2 \rfloor - 1} i \log(n/2) + \sum_{i=\lfloor n/2 \rfloor}^{n-1} i \log n \right]$$
(5.4)

Whether *n* is even or odd, one can show that the first sum of (5.4) does not exceed  $(n^2/8)\log(n/2)$ , which is  $(n^2/8)\log n - (n^2/8)$ , and the second sum does not exceed  $(3n^2/8)\log n$ . Thus, we can rewrite (5.4) as

$$P(n) \le 2 + \frac{8}{n^2} \left[ \frac{n^2}{2} \log n - \frac{n^2}{8} \right]$$
$$\le 1 + 4 \log n$$

as we wished to prove. This step completes the induction and shows that the average time to follow a path from the root to a random node of a binary search tree constructed by random insertions is  $O(\log n)$ , that is to within a constant factor as good as if the tree were complete. A more careful analysis shows that the constant 4 above is really about 1.4.

We can conclude from the above that the time of membership testing for a random member of the set takes  $O(\log n)$  time. A similar analysis shows that if we include in our average path length only those nodes that are missing both children, or only those missing left children, then the average path length still obeys an equation similar to (5.1), and is therefore  $O(\log n)$ . We may then conclude that testing membership of a random element not in the set, inserting a random new element, and deleting a random element also all take  $O(\log n)$  time on the average.

## **Evaluation of Binary Search Tree Performance**

Hash table implementations of dictionaries require constant time per operation on the average. Although this performance is better than that for a binary search tree, a hash table requires O(n) steps for the MIN operation, so if MIN is used frequently, the binary search tree will be the better choice; if MIN is not needed, we would probably prefer the hash table.

The binary search tree should also be compared with the partially ordered tree used for priority queues in Chapter 4. A partially ordered tree with n

#### 5.3 TRIES

elements requires only  $O(\log n)$  steps for each INSERT and DELETEMIN operation not only on the average, but also in the worst case. Moreover, the actual constant of proportionality in front of the logn factor will be smaller for a partially ordered tree than for a binary search tree. However, the binary search tree permits general DELETE and MIN operations, as well as the combination DELETEMIN, while the partially ordered tree permits only the latter. Moreover, MEMBER requires O(n) steps on a partially ordered tree but only  $O(\log n)$  steps on a binary search tree. Thus, while the partially ordered tree is well suited to implementing priority queues, it cannot do as efficiently any of the additional operations that the binary search tree can do.

#### 5.3 Tries

In this section we shall present a special structure for representing sets of character strings. The same method works for representing data types that are strings of objects of any type, such as strings of integers. This structure is known as the *trie*, derived from the middle letters of the word "retrieval."<sup>†</sup> By way of introduction, consider the following use of a set of character strings.

**Example 5.2.** As indicated in Chapter 1, one way to implement a spelling checker is to read a text file, break it into words (character strings separated by blanks and new lines), and find those words not in a standard dictionary of words in common use. Words in the text but not in the dictionary are printed out as possible misspellings. In Fig. 5.8 we see a sketch of one possible program *spell*. It makes use of a procedure getword(x, f) that sets x to be the next word in text file f; variable x is of a type called wordtype, which we shall define later. The variable A is of type SET; the SET operations we shall need are INSERT, DELETE, MAKENULL, and PRINT. The PRINT operator prints the members of the set.  $\Box$ 

The trie structure supports these set operations when the elements of the set are words, i.e., character strings. It is appropriate when many words begin with the same sequences of letters, that is, when the number of distinct prefixes among all the words in the set is much less than the total length of all the words.

In a trie, each path from the root to a leaf corresponds to one word in the represented set. This way, the nodes of the trie correspond to the prefixes of words in the set. To avoid confusion between words like THE and THEN, let us add a special *endmarker* symbol, \$, to the ends of all words, so no prefix of a word can be a word itself.

**Example 5.3.** In Fig. 5.9 we see a trie representing the set of words {THE, THEN, THIN, THIS, TIN, SIN, SING}. That is, the root corresponds to the empty string, and its two children correspond to the prefixes T and S. The

<sup>&</sup>lt;sup>†</sup> Trie was originally intended to be a homonym of "tree" but to distinguish these two terms many people prefer to pronounce trie as though it rhymes with "pie."

program spell ( input, output, dictionary ); type wordtype = { to be defined };  $SET = \{ to be defined, using the trie structure \};$ var A: SET; { holds input words not yet found in the dictionary } nextword: wordtype; dictionary: file of char; procedure getword ( var x: wordtype; f: file of char ); { a procedure to be defined that sets xto be the next word in file f } procedure INSERT ( x: wordtype; var S: SET ); { to be defined } procedure DELETE (x: wordtype; var S: SET ); { to be defined } procedure MAKENULL ( var S: SET ); { to be defined } procedure PRINT ( var S: SET ); { to be defined } begin MAKENULL(A); while not eof (input) do begin getword (nextword, input); INSERT(nextword, A) end: while not eof (dictionary) do begin getword (nextword, dictionary); DELETE(nextword, A) end; PRINT(A)end; { spell } Fig. 5.8. Sketch of spelling checker.

leftmost leaf represents the word THE, the next leaf the word THEN, and so on.  $\square$ 

We can make the following observations about the trie in Fig. 5.9.1. Each node has at most 27 children, one for each letter and \$.





Fig. 5.9. A trie.

- 2. Most nodes will have many fewer than 27 children.
- 3. A leaf reached by an edge labeled \$ cannot have any children, and may as well not be there.

#### **Trie Nodes as ADT's**

We can view a node of a trie as a mapping whose domain is  $\{A, B, \ldots, Z, \$\}$  (or whatever alphabet we choose) and whose value set is the type "pointer to trie node." Moreover, the trie itself can be identified with its root, so the ADT's TRIE and TRIENODE have the same data type, although their operations are substantially different. On TRIENODE we need the following operations:

- 1. procedure ASSIGN(node, c, p) that assigns value p (a pointer to a node) to character c in node node,
- 2. function VALUEOF(*node*, c) that produces the value associated with character c in *node*,<sup>†</sup> and
- 3. procedure GETNEW(node, c) to make the value of node for character c be a pointer to a new node.

Technically, we also need a procedure MAKENULL(*node*) to make *node* be the null mapping. One simple implementation of trie nodes is an array *node* of pointers to nodes, with the index set being  $\{A, B, \ldots, Z, \$\}$ . That is, we define

† VALUEOF is a function version of COMPUTE in Section 2.5.

```
type
```

chars = ('A', 'B', ..., 'Z', '\$'); TRIENODE = array[chars] of t TRIENODE;

If node is a trie node, node [c] is VALUEOF(node, c) for any c in the set chars. To avoid creating many leaves that are children corresponding to '\$', we shall adopt the convention that node ['\$'] is either nil or a pointer to the node itself. In the former case, node has no child corresponding to '\$', and in the latter case it is deemed to have such a child, although we never create it. Then we can write the procedures for trie nodes as in Fig. 5.10.

```
procedure MAKENULL ( var node: TRIENODE );
   { makes node a leaf, i.e., a null mapping }
   var
       c: chars:
   begin
       for c := 'A' to '$' do
          node[c] := nil
   end; { MAKENULL }
   procedure ASSIGN ( var node: TRIENODE; c: chars; p: † TRIENODE );
       begin
           node[c] := p
       end; { ASSIGN }
   function VALUEOF ( var node: TRIENODE; c: chars ) : † TRIENODE;
       begin
           return (node [c])
       end; { VALUEOF }
   procedure GETNEW ( var node: TRIENODE; c: chars );
       begin
           new(node[c]);
           MAKENULL(node [c])
       end; { GETNEW }
```

Fig. 5.10. Operations on trie nodes.

Now let us define

type TRIE = + TRIENODE;

We shall assume wordtype is an array of characters of some fixed length. The

5.3 TRIES

value of such an array will always be assumed to have at least one ''; we take the end of the represented word to be the first '', no matter what follows (presumably more '''s). On this assumption, we can write the procedure INSERT(*x*, words) to insert *x* into set words represented by a trie, as shown in Fig. 5.11. We leave the writing of MAKENULL, DELETE, and PRINT for tries represented as arrays for exercises.

```
procedure INSERT ( x: wordtype; var words: TRIE );
    var
         i: integer; { counts positions in word x }
         t: TRIE; { used to point to trie nodes
             corresponding to prefixes of x }
    begin
         i := 1;
         t := words;
         while x[i] <> '$' do begin
             if VALUEOF(t<sup>t</sup>, x[i]) = nil then
                 { if current node has no child for character x[i],
                      create one }
                 GETNEW(t^{\dagger}, x[i]);
             t := VALUEOF(t^{\dagger}, x[i]);
                  { proceed to the child of t for character x[i],
                      whether or not that child was just created }
             i := i+1 \{ \text{move along the word } x \}
         end;
         { now we have reached the first '$' in x }
         ASSIGN(tt, '$', t)
              { make loop for '$' to represent a leaf }
    end; {INSERT}
```

Fig. 5.11. The procedure INSERT.

#### A List Representation for Trie Nodes

The array representation of trie nodes takes a collection of words, having among them p different prefixes, and represents them with 27p bytes of storage. That amount of space could far exceed the total length of the words in the set. However, there is another implementation of tries that may save space. Recall that each trie node is a mapping, as discussed in Section 2.6. In principle, any implementation of mappings would do. Yet in practice, we want a representation suitable for mappings with a small domain and for mappings defined for comparatively few members of that domain. The linked list representation for mappings satisfies these requirements nicely. We may represent the mapping that is a trie node by a linked list of the characters for which the associated value is not the **nil** pointer. That is, a trie node is a linked list of cells of the type

```
type
```

```
celltype = record
  domain: chars;
  value: t celltype;
      { pointer to first cell on list for the child node }
    next: t celltype
      { pointer to next cell on the list }
end;
```

We shall leave the procedures ASSIGN, VALUEOF, MAKENULL, and GETNEW for this implementation of trie nodes as exercises. After writing these procedures, the INSERT operations on tries, in Fig. 5.11, and the other operations on tries that we left as exercises, should work correctly.

#### **Evaluation of the Trie Data Structure**

Let us compare the time and space needed to represent n words with a total of p different prefixes and a total length of l using a hash table and a trie. In what follows, we shall assume that pointers require four bytes. Perhaps the most space-efficient way to store words and still support INSERT and DELETE efficiently is in a hash table. If the words are of varying length, the cells of the buckets should not contain the words themselves; rather, the cells consist of two pointers, one to link the cells of the bucket and the other pointing to the beginning of a word belonging in the bucket.

The words themselves are stored in a large character array, and the end of each word is indicated by an endmarker character such as '\$'. For example, the words THE, THEN, and THIN could be stored as

#### THE\$THEN\$THIN\$ . . .

The pointers for the three words are cursors to positions 1, 5, and 10 of the array. The amount of space used in the buckets and character array is

- 1. 8n bytes for the cells of the buckets, there being one cell for each of the n words, and a cell has two pointers or 8 bytes,
- 2. l + n bytes for the character array to store the *n* words of total length l and their endmarkers.

The total space is thus 9n + l bytes plus whatever amount is used for the bucket headers.

In comparison, a trie with nodes implemented by linked lists requires p + n cells, one cell for each prefix and one cell for the end of each word. Each trie cell has a character and two pointers, and needs nine bytes, for a total space of 9n + 9p. If *l* plus the space for the bucket headers exceeds 9p, the trie uses less space. However, for applications such as storing a dictionary where l/p is typically less than 3, the hash table would use less space.

#### 5.4 BALANCED TREE IMPLEMENTATION OF SETS

In favor of the trie, however, let us note that we can travel down a trie, and thus perform operations INSERT, DELETE, and MEMBER in time proportional to the length of the word involved. A hash function to be truly "random" should involve each character of the word being hashed. It is fair, therefore, to state that computing the hash function takes roughly as much time as performing an operation like MEMBER on the trie. Of course the time spent computing the hash function does not include the time spent resolving collisions or performing the insertion, deletion, or membership test on the hash table, so we can expect tries to be considerably faster than hash tables for dictionaries whose elements are character strings.

Another advantage to the trie is that it supports the MIN operation efficiently, while hash tables do not. Further, in the hash table organization described above, we cannot easily reuse the space in the character array when a word is deleted (but see Chapter 12 for methods of handling such a problem).

#### 5.4 Balanced Tree Implementations of Sets

In Sections 5.1 and 5.2 we saw how sets could be implemented by binary search trees, and we saw that operations like INSERT could be performed in time proportional to the average depth of nodes in that tree. Further, we discovered that this average depth is  $O(\log n)$  for a "random" tree of n nodes. However, some sequences of insertions and deletions can produce binary search trees whose average depth is proportional to n. This suggests that we might try to rearrange the tree after each insertion and deletion so that it is always complete; then the time for INSERT and similar operations would always be  $O(\log n)$ .

In Fig. 5.12(a) we see a tree of six nodes that becomes the complete tree of seven nodes in Fig. 5.12(b), when element 1 is inserted. Every element in Fig. 5.12(a), however, has a different parent in Fig. 5.12(b), so we must take n steps to insert 1 into a tree like Fig. 5.12(a), if we wish to keep the tree as balanced as possible. It is thus unlikely that simply insisting that the binary search tree be complete will lead to an implementation of a dictionary, priority queue, or other ADT that includes INSERT among its operations, in  $O(\log n)$  time.

There are several other approaches that do yield worst case  $O(\log n)$  time per operation for dictionaries and priority queues, and we shall consider one of them, called a "2-3 tree," in detail. A 2-3 tree is a tree with the following two properties.

1. Each interior node has two or three children.

2. Each path from the root to a leaf has the same length.

We shall also consider a tree with zero nodes or one node as special cases of a 2-3 tree.

We represent sets of elements that are ordered by some linear order <, as follows. Elements are placed at the leaves; if element a is to the left of



Fig. 5.12. Complete trees.

element b, then a < b must hold. We shall assume that the "<" ordering of elements is based on one field of a record that forms the element type; this field is called the *key*. For example, elements might represent people and certain information about people, and in that case, the key field might be "social security number."

At each interior node we record the key of the smallest element that is a descendant of the second child and, if there is a third child, we record the key of the smallest element descending from that child as well.<sup>†</sup> Figure 5.13 is an example of a 2-3 tree. In that and subsequent examples, we shall identify an element with its key field, so the order of elements becomes obvious.



Fig. 5.13. A 2-3 tree.

<sup>†</sup> There is another version of 2-3 trees that places whole records at interior nodes, as a binary search tree does.

5.4 BALANCED TREE IMPLEMENTATIONS OF SETS

Observe that a 2-3 tree of k levels has between  $2^{k-1}$  and  $3^{k-1}$  leaves. Put another way, a 2-3 tree representing a set of n elements requires at least  $1 + \log_3 n$  levels and no more than  $1 + \log_2 n$  levels. Thus, path lengths in the tree are  $O(\log n)$ .

We can test membership of a record with key x in a set represented by a 2-3 tree in  $O(\log n)$  time by simply moving down the tree, using the values of the elements recorded at the interior nodes to guide our path. At a node node, compare x with the value y that represents the smallest element descending from the second child of node. (Recall we are treating elements as if they consisted solely of a key field.) If x < y, move to the first child of node. If  $x \ge y$ , and node has only two children, move to the second child of node. If node has three children and  $x \ge y$ , compare x with z, the second value recorded at node, the value that indicates the smallest descendant of the third child of node. If x < z, go to the second child, and if  $x \ge z$ , go to the third child. In this manner, we find ourselves at a leaf eventually, and x is in the represented set if and only if x is at the leaf. Evidently, if during this process we find x = y or x = z, we can stop immediately. However, we stated the algorithm as we did because in some cases we shall wish to find the leaf with x as well as to verify its existence.

#### **Insertion into a 2-3 Tree**

To insert a new element x into a 2-3 tree, we proceed at first as if we were testing membership of x in the set. However, at the level just above the leaves, we shall be at a node *node* whose children, we discover, do not include x. If *node* has only two children, we simply make x the third child of *node*, placing the children in the proper order. We then adjust the two numbers at *node* to reflect the new situation.

For example, if we insert 18 into Fig. 5.13, we wind up with *node* equal to the rightmost node at the middle level. We place 18 among the children of *node*, whose proper order is 16, 18, 19. The two values recorded at *node* become 18 and 19, the elements at the second and third children. The result is shown in Fig. 5.14.

Suppose, however, that x is the fourth, rather than the third child of *node*. We cannot have a node with four children in a 2-3 tree, so we split *node* into two nodes, which we call *node* and *node'*. The two smallest elements among the four children of *node* stay with *node*, while the two larger elements become children of *node'*. Now, we must insert *node'* among the children of *p*, the parent of *node*. This part of the insertion is analogous to the insertion of a leaf as a child of *node*. That is, if *p* had two children, we make *node'* the third and place it immediately to the right of *node*. If *p* had three children before *node'* was created, we split *p* into *p* and *p'*, giving *p* the two leftmost children and *p'* the remaining two, and then we insert *p'* among the children of *p*'s parent, recursively.

One special case occurs when we wind up splitting the root. In that case we create a new root, whose two children are the two nodes into which the



Fig. 5.14. 2-3 tree with 18 inserted.

old root was split. This is how the number of levels in a 2-3 tree increases.

**Example 5.4.** Suppose we insert 10 into the tree of Fig. 5.14. The intended parent of 10 already has children 7, 8, and 12, so we split it into two nodes. The first of these has children 7 and 8; the second has 10 and 12. The result is shown in Fig. 5.15(a). We must now insert the new node with children 10 and 12 in its proper place as a child of the root of Fig. 5.15(a). Doing so gives the root four children, so we split it, and create a new root, as shown in Fig. 5.15(b). The details of how information regarding smallest elements of subtrees is carried up the tree will be given when we develop the program for the command INSERT.  $\Box$ 

### **Deletion in a 2-3 tree**

When we delete a leaf, we may leave its parent *node* with only one child. If *node* is the root, delete *node* and let its lone child be the new root. Otherwise, let p be the parent of *node*. If p has another child, adjacent to *node* on either the right or the left, and that child of p has three children, we can transfer the proper one of those three to *node*. Then *node* has two children, and we are done.

If the children of p adjacent to *node* have only two children, transfer the lone child of *node* to an adjacent sibling of *node*, and delete *node*. Should p now have only one child, repeat all the above, recursively, with p in place of *node*.

**Example 5.5.** Let us begin with the tree of Fig. 5.15(b). If 10 is deleted, its parent has only one child. But the grandparent has another child that has three children, 16, 18, and 19. This node is to the right of the deficient node, so we pass the deficient node the smallest element, 16, leaving the 2-3 tree in Fig. 5.16(a).



Fig. 5.15. Insertion of 10 into the tree of Fig. 5.14.

Next suppose we delete 7 from the tree of Fig. 5.16(a). Its parent now has only one child, 8, and the grandparent has no child with three children. We therefore make 8 be a sibling of 2 and 5, leaving the tree of Fig. 5.16(b). Now the node starred in Fig. 5.16(b) has only one child, and its parent has no other child with three children. Thus we delete the starred node, making its child be a child of the sibling of the starred node. Now the root has only one child, and we delete it, leaving the tree of Fig. 5.16(c).

Observe in the above examples, the frequent manipulation of the values at interior nodes. While we can always calculate these values by walking the tree, it can be done as we manipulate the tree itself, provided we remember the smallest value among the descendants of each node along the path from the root to the deleted leaf. This information can be computed by a recursive deletion algorithm, with the call at each node being passed, from above, the correct quantity (or the value "minus infinity" if we are along the leftmost path). The details require careful case analysis and will be sketched later when we consider the program for the DELETE operation.  $\Box$ 

#### **Data Types for 2-3 Trees**

Let us restrict ourselves to representing by 2-3 trees sets of elements whose keys are real numbers. The nature of other fields that go with the field key, to make up a record of type elementtype, we shall leave unspecified, as it has no bearing on what follows.



#### Fig. 5.16. Deletion in a 2-3 tree.

In Pascal, the parents of leaves must be records consisting of two reals (the keys of the smallest elements in the second and third subtrees) and of three pointers to elements. The parents of these nodes are records consisting of two reals and of three pointers to parents of leaves. This progression continues indefinitely; each level in a 2-3 tree is of a different type from all other levels. Such a situation would make programming 2-3 tree operations in Pascal impossible, but fortunately, Pascal provides a mechanism, the variant record structure, that enables us to regard all 2-3 tree nodes as having the same type, even though some are elements, and some are records with pointers and reals.<sup>†</sup> We can define nodes as in Fig. 5.17. Then we declare a set, represented by a 2-3 tree, to be a pointer to the root as in Fig. 5.17.

#### **Implementation of INSERT**

The details of operations on 2-3 trees are quite involved, although the principles are simple. We shall therefore describe only one operation, insertion, in detail; the others, deletion and membership testing, are similar in spirit, and finding the minimum is a trivial search down the leftmost path. We shall write the insertion routine as main procedure, INSERT, which we call at the root, and a procedure *insert*1, which gets called recursively down the tree.

<sup>&</sup>lt;sup>†</sup> All nodes, however, take the largest amount of space needed for any of the variant types, so Pascal is not really the best language for implementing 2-3 trees in practice.

5.4 BALANCED TREE IMPLEMENTATIONS OF SETS

```
type
    elementtype = record
        key: real;
        {other fields as warranted}
    end;
    nodetypes = (leaf, interior);
    twothreenode = record
        case kind: nodetypes of
        leaf: (element: elementtype);
        interior: (firstchild, secondchild, thirdchild: t twothreenode;
        lowofsecond, lowofthird: real)
end;
SET = t twothreenode;
```

Fig. 5.17. Definition of a node in a 2-3 tree.

For convenience, we assume that a 2-3 tree is not a single node or empty. These two cases require a straightforward sequence of steps which the reader is invited to provide as an exercise.

```
procedure insert1 ( node: † twothreenode;
```

x: elementtype; { x is to be inserted into the subtree of node }
var pnew: t twothreenode; { pointer to new node created to right of node }
var low: real ); { smallest element in the subtree pointed to by pnew }

begin

pnew := nil; if node is a leaf then begin if x is not the element at node then begin create new node pointed to by pnew; put x at the new node; low := x.keyend end else begin { node is an interior node } let w be the child of node to whose subtree x belongs; insert1(w, x, pback, lowback); if pback <> nil then begin insert pointer pback among the children of node just to the right of w; if node has four children then begin create new node pointed to by pnew; give the new node the third and fourth children of node; adjust lowofsecond and lowofthird in node and the new node;

set *low* to be the lowest key among the children of the new node

end end

end; { insert 1 }

Fig. 5.18. Sketch of 2-3 tree insertion program.

We would like *insert* 1 to return both a pointer to a new node, if it must create one, and the key of the smallest element descended from that new node. As the mechanism in Pascal for creating such a function is awkward, we shall instead declare *insert* 1 to be a procedure that assigns values to parameters *pnew* and *low* in the case that it must "return" a new node. We sketch *insert* 1 in Fig. 5.18. The complete procedure is shown in Fig. 5.19; some comments in Fig. 5.18 have been omitted from Fig. 5.19 to save space.

```
procedure insert 1 ( node: t twothreenode; x: elementtype;
var pnew: t twothreenode; var low: real );
```

var

pback: t twothreenode; lowback: real; child: 1..3; { indicates which child of node is followed in recursive call (cf. w in Fig. 5.18) } w: t twothreenode; { pointer to the child }

begin

```
pnew := nil;
if node t.kind = leaf then begin
    if node t.kind = leaf then begin
        { create new leaf holding x and "return" this node }
        new(pnew, leaf);
        if (node t.element.key < x.key) then
        { place x in new node to right of current node }
        begin pnew t.element := x; low := x.key end
        else begin { x belongs to left of element at current node }
            pnew t.element := node t.element;
            node t.element := x;
            low := pnew t.element.key
        end
        end
```

end

else begin { node is an interior node }

{ select the child of *node* that we must follow }

if x.key < node t.low of second then

begin child := 1; w := node t.firstchild end

```
5.4 BALANCED TREE IMPLEMENTATIONS OF SETS
                                                                            177
            else if (node t.thirdchild = nil) or (x.key < node t.lowofthird) then begin
                \{x \text{ is in second subtree}\}
                child := 2;
                w := node t.secondchild
            end
            else begin { x is in third subtree }
                child := 3:
                w := node t.thirdchild
            end;
            insert 1(w, x, pback, lowback);
            if pback <> nil then
                { a new child of node must be inserted }
                if node t.thirdchild = nil then
                    { node had only two children, so insert new node in proper place }
                    if child = 2 then begin
                         node t.thirdchild := pback;
                         node t.lowofthird := lowback
                    end
                     else begin { child = 1 }
                         node t.thirdchild := node t.secondchild;
                         node t.lowofthird := node t.lowofsecond;
                         node t.secondchild := pback;
                         node t.lowofsecond := lowback
                    end
                else begin { node already had three children }
                    new (pnew, interior);
                    if child = 3 then begin
                        { pback and third child become children of new node }
                        pnew t.firstchild := node t.thirdchild;
                        pnew i.secondchild := pback;
                        pnew t.thirdchild := nil;
                        pnew t.lowofsecond := lowback;
                             { lowofthird is undefined for pnew }
                         low := node t.lowofthird;
                        node t.thirdchild := nil
                    end
                    else begin { child \leq 2; move third child of node to pnew}
                        pnew t.secondchild := node t.thirdchild;
                        pnew t.lowofsecond := node t.lowofthird;
                        pnew t.thirdchild := nil;
                        node t.thirdchild := nil
                    end;
                    if child = 2 then begin
                        { pback becomes first child of pnew }
                        pnew t.firstchild := pback;
```

```
low := lowback
end;
if child = 1 then begin
{ second child of node is moved to pnew;
        pback becomes second child of node }
        pnew t.firstchild := node t.secondchild;
        low := node t.lowofsecond;
        node t.secondchild := pback;
        node t.lowofsecond := lowback
end
```

end

end end; { insert 1 }

#### Fig. 5.19. The procedure insert 1.

Now we can write the procedure INSERT, which calls *insert* 1. If *insert* 1 "returns" a new node, then INSERT must create a new root. The code is shown in Fig. 5.20 on the assumption that the type SET is t twothreenode, i.e., a pointer to the root of a 2-3 tree whose leaves contain the members of the set.

```
procedure INSERT ( x: elementtype; var S: SET );
```

var

pback: t twothreenode; { pointer to new node returned by insert 1 }
lowback: real; { low value in subtree of pback }

saveS: SET; { place to store a temporary copy of the pointer S } begin

{ checks for S being empty or a single node should occur here,

and an appropriate insertion procedure should be included }

insert 1(S, x, pback, lowback);

if pback <> nil then begin

{ create new root; its children are now pointed to by S and pback } saveS := S;

```
new(S);
```

St.firstchild := saveS;

St.secondchild := pback;

St.lowofsecond := lowback;

 $S \uparrow$ .thirdchild := nil;

end

end; { INSERT }

Fig. 5.20. INSERT for sets represented by 2-3 trees.

#### 5.4 BALANCED TREE IMPLEMENTATIONS OF SETS

#### **Implementation of DELETE**

We shall sketch a function delete 1 that takes a pointer to a node node and an element x, and deletes a leaf descended from node having value x, if there is one.<sup>†</sup> Function delete 1 returns true if after deletion node has only one child, and it returns false if node still has two or three children. A sketch of the code for delete 1 is shown in Fig. 5.21.

We leave the detailed code for function *delete* 1 for the reader. Another exercise is to write a procedure DELETE(S, x) that checks for the special cases that the set S consists only of a single leaf or is empty, and otherwise calls *delete* 1(S, x); if *delete* 1 returns true, the procedure removes the root (the node pointed to by S) and makes S point to its lone child.

function delete 1 ( node: t twothreenode; x: elementtype ) : boolean;
var

orlyone: boolean; { to hold the value returned by a call to delete 1 } begin

delete 1 := false;

if the children of node are leaves then begin

if x is among those leaves then begin

remove x;

shift children of *node* to the right of x one position left; if *node* now has one child **then** 

delete 1 := true

end

end

else begin { node is at level two or higher }

determine which child of node could have x as a descendant;  $onlyone := delete 1(w, x); \{w \text{ stands for node t.firstchild,} \}$ 

node t.secondchild, or node t.thirdchild, as appropriate }

if onlyone then begin { fix children of node }

if w is the first child of node then

- if y, the second child of node, has three children then make the first child of y be the second child of w else begin { y has two children }
- make the child of w be the first child of y;

remove w from among the children of node;

if node now has one child then

delete 1 := true

end;

if w is the second child of node then

if y, the first child of *node*, has three children then make the third child of y be the first child of w

† A useful variant would take only a key value and delete any element with that key.

else { y has two children }

if z, the third child of node, exists and has three children then

end;

if w is the third child of node then

if y, the second child of node, has three children then make the third child of y be the first child of w else begin { y has two children }

make the child of w be the third child of y;

remove w from among the children of node

end { note node surely has two children left in this case }

end end

end; { delete 1 }

Fig. 5.21. Recursive deletion procedure.

#### 5.5 Sets with the MERGE and FIND Operations

In certain problems we start with a collection of objects, each in a set by itself; we then combine sets in some order, and from time to time ask which set a particular object is in. These problems can be solved using the operations MERGE and FIND. The operation MERGE(A, B, C) makes C equal to the union of sets A and B, provided A and B are disjoint (have no member in common); MERGE is undefined if A and B are not disjoint. FIND(x) is a function that returns the set of which x is a member; in case x is in two or more sets, or in no set, FIND is not defined.

**Example 5.6.** An *equivalence relation* is a reflexive, symmetric, and transitive relation. That is, if  $\equiv$  is an equivalence relation on set S, then for any (not necessarily distinct) members a, b, and c in S, the following properties hold:

- 1.  $a \equiv a$  (reflexivity)
- 2. If  $a \equiv b$ , then  $b \equiv a$  (symmetry).
- 3. If  $a \equiv b$  and  $b \equiv c$ , then  $a \equiv c$  (transitivity).

The relation "is equal to" (=) is the paradigm equivalence relation on any set S. For a, b, and c in S, we have (1) a = a, (2) if a = b, then b = a, and (3) if a = b and b = c, then a = c. There are many other equivalence relations, however, and we shall shortly see several additional examples.

#### 5.5 SETS WITH THE MERGE AND FIND OPERATIONS

In general, whenever we partition a collection of objects into disjoint groups, the relation  $a \equiv b$  if and only if a and b are in the same group is an equivalence relation. "Is equal to" is the special case where every element is in a group by itself.

More formally, if a set S has an equivalence relation defined on it, then the set S can be partitioned into disjoint subsets  $S_1, S_2, \ldots$ , called equivalence classes, whose union is S. Each subset  $S_i$  consists of equivalent members of S. That is,  $a \equiv b$  for all a and b in  $S_i$ , and  $a \neq b$  if a and b are in different subsets. For example, the relation congruence modulo  $n^{\dagger}$  is an equivalence relation on the set of integers. To check that this is so, note that a-a = 0, which is a multiple of n (reflexivity); if a-b = dn, then b-a = (-d)n (symmetry); and if a-b = dn and b-c = en, then a-c = (d+e)n (transitivity). In the case of congruence modulo n there are n equivalence classes, which are the set of integers congruent to 0, the set of integers congruent to 1, ..., the set of integers congruent to n-1.

The equivalence problem can be formulated in the following manner. We are given a set S and a sequence of statements of the form "a is equivalent to b." We are to process the statements in order in such a way that at any time we are able to determine in which equivalence class a given element belongs. For example, suppose  $S = \{1, 2, ..., 7\}$  and we are given the sequence of statements

#### 1=2 5=6 3=4 1=4

to process. The following sequence of equivalence classes needs to be constructed, assuming that initially each element of S is in an equivalence class by itself.

1≡2	$\{1,2\}\ \{3\}\ \{4\}\ \{5\}\ \{6\}\ \{7\}$
5≡6	$\{1,2\}$ $\{3\}$ $\{4\}$ $\{5,6\}$ $\{7\}$
3≡4	$\{1,2\}\ \{3,4\}\ \{5,6\}\ \{7\}$
1≡4	{1,2,3,4} {5,6} {7}

We can "solve" the equivalence problem by starting with each element in a named set. When we process statement a=b, we FIND the equivalence classes of a and b and then MERGE them. We can at any time use FIND to tell us the current equivalence class of any element.

The equivalence problem arises in several areas of computer science. For example, one form occurs when a Fortran compiler has to process "equivalence declarations" such as

<sup>†</sup> We say a is congruent to b modulo n if a and b have the same remainders when divided by n, or put another way, a-b is a multiple of n.

### EQUIVALENCE (A(1),B(1,2),C(3)), (A(2),D,E), (F,G)

Another example, presented in Chapter 7, uses solutions to the equivalence problem to help find minimum-cost spanning trees.  $\Box$ 

#### A Simple Implementation of MFSET

Let us begin with a simplified version of the MERGE-FIND ADT. We shall define an ADT, called MFSET, consisting of a set of subsets, which we shall call *components*, together with the following operations:

- 1. MERGE(A, B) takes the union of the components A and B and calls the result either A or B, arbitrarily.
- 2. FIND(x) is a function that returns the name of the component of which x is a member.
- 3. INITIAL(A, x) creates a component named A that contains only the element x.

To make a reasonable implementation of MFSET, we must restrict our underlying types, or alternatively, we should recognize that MFSET really has two other types as "parameters" — the type of set names and the type of members of these sets. In many applications we can use integers as set names. If we take n to be the number of elements, we may also use integers in the range [1..n] for the members of components. For the implementation we have in mind, it is important that the type of set members be a subrange type, because we want to index into an array defined over that subrange. The type of set names is not important, as this type is the type of array elements, not their indices. Observe, however, that if we wanted the member type to be other than a subrange type, we could create a mapping, with a hash table, for example, that assigned these to unique integers in a subrange. We only need to know the total number of elements in advance.

The implementation we have in mind is to declare

const

 $n = \{ \text{ number of elements } \};$ type

MFSET = array[1..n] of integer;

as a special case of the more general type

#### array[subrange of members] of (type of set names);

Suppose we declare *components* to be of type MFSET with the intention that *components* [x] holds the name of the set currently containing x. Then the three MFSET operations are easy to write. For example, the operation MERGE is shown in Fig. 5.22. INITIAL(A, x) simply sets *components* [x] to A, and FIND(x) returns *components* [x].

The time performance of this implementation of MFSET is easy to

5.5 SETS WITH THE MERGE AND FIND OPERATIONS

procedure MERGE ( A, B: integer; var C: MFSET );
 var
 x: 1..n;
 begin
 for x := 1 to n do
 if C[x] = B then
 C[x] := A
end; { MERGE }

Fig. 5.22. The procedure MERGE.

analyze. Each execution of the procedure MERGE takes O(n) time. On the other hand, the obvious implementations of INITIAL(A, x) and FIND(x) have constant running times.

#### **A Faster Implementation of MFSET**

Using the algorithm in Fig. 5.22, a sequence of n-1 MERGE instructions will take  $O(n^2)$  time.<sup>†</sup> One way to speed up the MERGE operation is to link together all members of a component in a list. Then, instead of scanning all members when we merge component *B* into *A*, we need only run down the list of members of *B*. This arrangement saves time on the average. However, it could happen that the *i*<sup>th</sup> merge is of the form MERGE(*A*, *B*) where *A* is a component of size 1 and *B* is a component of size *i*, and that the result is named *A*. This merge operation would require O(i) steps, and a sequence of n-1 such merge instructions would take on the order of  $\sum_{i=1}^{n-1} i = n(n-1)/2$  time.

One way to avoid this worst case situation is to keep track of the size of each component and always merge the smaller into the larger.<sup>‡</sup> Thus, every time a member is merged into a bigger component, it finds itself in a component at least twice as big. Thus, if there are initially n components, each with one member, none of the n members can have its component changed more than  $1+\log n$  times. As the time spent by this new version of MERGE is proportional to the number of members whose component names are changed, and the total number of such changes is at most  $n(1+\log n)$ , we see that  $O(n \log n)$  work suffices for all merges.

Now let us consider the data structure needed for this implementation. First, we need a mapping from set names to records consisting of

 $<sup>\</sup>dagger$  Note that n-1 is the largest number of merges that can be performed before all elements are in one set.

<sup>&</sup>lt;sup>‡</sup> Note that our ability to call the resulting component by the name of either of its constituents is important here, although in the simpler implementation, the name of the first argument was always picked.

1. a count giving the number of members in the set and

2. the index in the array of the first element of that set.

We also need another array of records, indexed by members, to indicate

1. the set of which each element is a member and

2. the next array element on the list for that set.

We use 0 to serve as NIL, the end-of-list marker. In a language that lent itself to such constructs, we would prefer to use pointers in this array, but Pascal does not permit pointers into arrays.

In the special case where set names, as well as members, are chosen from the subrange 1..n, we can use an array for the mapping described above. That is, we define

```
type
```

```
nametype = 1..n;
elementtype = 1..n;
MFSET = record
setheaders: array[1..n] of record
{ headers for set lists }
count: 0..n;
firstelement: 0..n
end;
names: array[1..n] of record
{ table giving set containing each member }
setname: nametype;
nextelement: 0..n
end
end;
```

The procedures INITIAL, MERGE, and FIND are shown in Fig. 5.23.

Figure 5.24 shows an example of the data structure used in Fig. 5.23, where set 1 is  $\{1, 3, 4\}$ , set 2 is  $\{2\}$ , and set 5 is  $\{5, 6\}$ .

#### A Tree Implementation of MFSET's

Another, completely different, approach to the implementation of MFSET's uses trees with pointers to parents. We shall describe this approach informally. The basic idea is that nodes of trees correspond to set members, with an array or other implementation of a mapping leading from set members to their nodes. Each node, except the root of each tree, has a pointer to its parent. The roots hold the name of the set, as well as an element. A mapping from set names to roots allows access to any given set, when merges are done.

Figure 5.25 shows the sets  $A = \{1, 2, 3, 4\}$ ,  $B = \{5, 6\}$ , and  $C = \{7\}$  represented in this form. The rectangles are assumed to be part of the root node, not separate nodes.

5.5 SETS WITH THE MERGE AND FIND OPERATIONS

```
procedure INITIAL ( A: nametype; x: elementtype; var C: MFSET );
    { initialize A to a set containing x only }
    begin
        C.names[x].setname := A;
        C.names[x].nextelement := 0;
             { null pointer at end of list of members of A }
        C.setheaders [A].count := 1;
        C.setheaders [A].firstelement := x
    end; { INITIAL }
procedure MERGE (A, B: nametype; var C: MFSET );
    { merge A and B, calling the result A or B, arbitrarily }
    var
        i: 0..n; { used to find end of smaller list }
    begin
        if C.setheaders [A].count > C.setheaders [B].count then begin
             { A is the larger set; merge B into A }
            { find end of B, changing set names to A as we go }
            i := C.setheaders[B].firstelement;
            while C.names [i].nextelement <> 0 do begin
                 C.names[i].setname := A;
                i := C.names [i].nextelement
            end:
            { append list A to the end of B and call the result A }
            { now i is the index of the last member of B }
            C.names [i].setname := A;
            C.names [i].nextelement := C.setheaders [A].firstelement;
            C.setheaders [A].firstelement := C.setheaders [B].firstelement;
            C.setheaders [A].count := C.setheaders [A].count +
                         C.setheaders [B].count;
            C.setheaders [B].count := 0;
            C.setheaders [B].firstelement := 0
                { above two steps not really necessary, as set B no longer exists }
        end
        else { B is at least as large as A }
            { code similar to case above, but with A and B interchanged }
    end; { MERGE }
function FIND (x: 1...n; var C: MFSET );
    { return the name of the set of which x is a member }
    begin
        return (C.names [x].setname)
   end; { FIND }
```

Fig. 5.23. The operations of an MFSET.

To find the set containing an element x, we first consult a mapping (e.g., an array) not shown in Fig. 5.25, to obtain a pointer to the node for x. We then follow the path from that node to the root of its tree and read the name of the set there.

The basic merge operation is to make the root of one tree be a child of the root of the other. For example, we could merge A and B of Fig. 5.25 and call the result A, by making node 5 a child of node 1. The result is shown in Fig. 5.26. However, indiscriminate merging could result in a tree of n nodes that is a single chain. Then doing a FIND operation on each of those nodes would take  $O(n^2)$  time. Observe that although a merge can be done in O(1)steps, the cost of a reasonable number of FIND's will dominate the total cost, and this approach is not necessarily better than the simplest one for executing n merges and n finds.

However, a simple improvement guarantees that if n is the number of elements, then no FIND will take more than  $O(\log n)$  steps. We simply keep at each root a count of the number of elements in the set, and when called upon to merge two sets, we make the root of the smaller tree be a child of the root of the larger. Thus, every time a node is moved to a new tree, two things happen: the distance from the node to its root increases by one, and the node will be in a set with at least twice as many elements as before. Thus, if n is the total number of elements, no node can be moved more than  $\log n$  times; hence, the distance to its root can never exceed  $\log n$ . We conclude that each FIND requires at most  $O(\log n)$  time.

#### **Path Compression**

Another idea that may speed up this implementation of MFSET's is *path* compression. During a FIND, when following a path from some node to the root, make each node encountered along the path be a child of the root. The easiest way to do this is in two passes. First, find the root, and then retraverse the same path, making each node a child of the root.

**Example 5.7.** Figure 5.27(a) shows a tree before executing a FIND operation on the node for element 7 and Fig. 5.27(b) shows the result after 5 and 7 are made children of the root. Nodes 1 and 2 on the path are not moved because 1 is the root, and 2 is already a child of the root.  $\Box$ 

Path compression does not affect the cost of MERGE's; each MERGE still takes a constant amount of time. There is, however, a subtle speedup in FIND's since path compression tends to shorten a large number of paths from various nodes to the root with relatively little effort.

Unfortunately, it is very difficult to analyze the average cost of FIND's when path compression is used. It turns out that if we do not require that smaller trees be merged into larger ones, we require no more than  $O(n\log n)$  time to do *n* FIND's. Of course, the first FIND may take O(n) time by itself for a tree consisting of one chain. But path compression can change a tree very rapidly and no matter in what order we apply FIND to elements of any tree no more than O(n) time is spent on *n* FIND's. However, there are



Fig. 5.24. Example of the MFSET data structure.



Fig. 5.25. MFSET represented by a collection of trees.

sequences of MERGE and FIND instructions that require  $\Omega(n \log n)$  time.

The algorithm that both uses path compression and merges the smaller tree into the larger is asymptotically the most efficient method known for implementing MFSET's. In particular, n FIND's require no more than  $O(n\alpha(n))$  time, where  $\alpha(n)$  is a function that is not constant, yet grows much more slowly than log n. We shall define  $\alpha(n)$  below, but the analysis that leads to this bound is beyond the scope of this book.






#### 5.6 AN ADT WITH MERGE AND SPLIT

# The Function $\alpha(n)$

The function  $\alpha(n)$  is closely related to a very rapidly growing function A(x, y), known as Ackermann's function. A(x, y) is defined recursively by:

$$A(0, y) = 1 \text{ for } y \ge 0$$
  

$$A(1, 0) = 2$$
  

$$A(x, 0) = x+2 \text{ for } x \ge 2$$
  

$$A(x, y) = A(A(x-1, y), y-1) \text{ for } x, y \ge 1$$

Each value of y defines a function of one variable. For example, the third line above tells us that for y=0, this function is "add 2." For y = 1, we have A(x, 1) = A(A(x-1, 1), 0) = A(x-1, 1) + 2, for x > 1, with A(1, 1) = A(A(0, 1), 0) = A(1, 0) = 2. Thus A(x, 1) = 2x for all  $x \ge 1$ . In other words, A(x, 1) is "multiply by 2." Then, A(x, 2) = A(A(x-1, 2), 1) = 2A(x-1, 2) for x > 1. Also,  $A(1, 2) = A(A(0, 2), 1) = A(A(x-1, 2), 1) = A(x, 2) = 2^x$ . Similarly, we can show that  $A(x, 3) = 2^{2^{-1}}$  (stack of x 2's), while A(x, 4) is so rapidly growing there is no accepted mathematical notation for such a function.

A single-variable Ackermann's function can be defined by letting A(x) = A(x, x). The function  $\alpha(n)$  is a pseudo-inverse of this single variable function. That is,  $\alpha(n)$  is the least x such that  $n \le A(x)$ . For example, A(1) = 2, so  $\alpha(1) = \alpha(2) = 1$ . A(2) = 4, so  $\alpha(3) = \alpha(4) = 2$ . A(3) = 8, so  $\alpha(5) = \cdots = \alpha(8) = 3$ . So far,  $\alpha(n)$  seems to be growing rather steadily.

However, A(4) is a stack of 65536 2's. Since  $\log(A(4))$  is a stack of 65535 2's, we cannot hope even to write uA(4) explicitly, as it would take  $\log(A(4))$  bits to do so. Thus  $\alpha(n) \leq 4$  for all integers *n* one is ever likely to encounter. Nevertheless,  $\alpha(n)$  eventually reaches 5, 6, 7, . . . on its unimaginably slow course toward infinity.

## 5.6 An ADT with MERGE and SPLIT

Let S be a set whose members are ordered by the relation <. The operation SPLIT(S,  $S_1$ ,  $S_2$ , x) partitions S into two sets:  $S_1 = \{a \mid a \text{ is in } S \text{ and } a < x\}$ and  $S_2 = \{a \mid a \text{ is in } S \text{ and } a \ge x\}$ . The value of S after the split is undefined, unless it is one of  $S_1$  or  $S_2$ . There are several situations where the operation of splitting sets by comparing each member with a fixed value x is essential. We shall consider one such problem here.

#### The Longest Common Subsequence Problem

A subsequence of a sequence x is obtained by removing zero or more (not necessarily contiguous) elements from x. Given two sequences x and y, a longest common subsequence (LCS) is a longest sequence that is a subsequence of both x and y.

For example, an LCS of 1, 2, 3, 2, 4, 1, 2 and 2, 4, 3, 1, 2, 1 is the subsequence 2, 3, 2, 1, formed as shown in Fig. 5.28. There are other LCS's as well, such as 2, 4, 1, 2, but there are no common subsequences of length 5.



Fig. 5.28. A longest common subsequence.

There is a UNIX command called *diff* that compares files line-by-line, finding a longest common subsequence, where a line of a file is considered an element of the subsequence. That is, whole lines are analogous to the integers 1, 2, 3, and 4 in Fig. 5.28. The assumption behind the command *diff* is that the lines of each file that are not in this LCS are lines inserted, deleted or modified in going from one file to the other. For example, if the two files are versions of the same program made several days apart, *diff* will, with high probability, find the changes.

There are several general solutions to the LCS problem that work in  $O(n^2)$  steps on sequences of length n. The command *diff* uses a different strategy that works well when the files do not have too many repetitions of any line. For example, programs will tend to have lines "begin" and "end" repeated many times, but other lines are not likely to repeat.

The algorithm used by diff for finding an LCS makes use of an efficient implementation of sets with operations MERGE and SPLIT, to work in time  $O(p \log n)$ , where n is the maximum number of lines in a file and p is the number of pairs of positions, one from each file, that have the same line. For example, p for the strings in Fig. 5.28 is 12. The two 1's in each string contribute four pairs, the 2's contribute six pairs, and 3 and 4 contribute one pair each. In the worst case, p could be  $n^2$ , and this algorithm would take  $O(n^2 \log n)$  time. However, in practice, p is usually closer to n, so we can expect an  $O(n \log n)$  time complexity.

To begin the description of the algorithm let  $A = a_1a_2 \cdots a_n$  and  $B = b_1b_2 \cdots b_m$  be the two strings whose LCS we desire. The first step is to tabulate for each value a, the positions of the string A at which a appears. That is, we define PLACES(a) = {  $i \mid a = a_i$  }. We can compute the sets PLACES(a) by constructing a mapping from symbols to headers of lists of positions. By using a hash table, we can create the sets PLACES(a) in O(n) "steps" on the average, where a "step" is the time it takes to operate on a symbol, say to hash it or compare it with another. This time could be a constant if symbols are characters or integers, say. However, if the symbols of A

#### 5.6 AN ADT WITH MERGE AND SPLIT

and B are really lines of text, then steps take an amount of time that depends on the average length of a line of text.

Having computed PLACES(a) for each symbol a that occurs in string A, we are ready to find an LCS. To simplify matters, we shall only show how to find the length of the LCS, leaving the actual construction of the LCS as an exercise. The algorithm considers each  $b_j$ , for  $j = 1, 2, \ldots, m$ , in turn. After considering  $b_j$ , we need to know, for each i between 0 and n, the length of the LCS of strings  $a_1 \cdots a_i$  and  $b_1 \cdots b_j$ .

We shall group values of *i* into sets  $S_k$ , for k = 0, 1, ..., n, where  $S_k$  consists of all those integers *i* such that the LCS of  $a_1 \cdots a_i$  and  $b_1 \cdots b_j$  has length *k*. Note that  $S_k$  will always be a set of consecutive integers, and the integers in  $S_{k+1}$  are larger than those in  $S_k$ , for all *k*.

**Example 5.8.** Consider Fig. 5.28, with j = 5. If we try to match zero symbols from the first string with the first five symbols of the second (24312), we naturally have an LCS of length 0, so 0 is in  $S_0$ . If we use the first symbol from the first string, we can obtain an LCS of length 1, and if we use the first two symbols, 12, we can obtain an LCS of length 2. However, using 123, the first three symbols, still gives us an LCS of length 2 when matched against 24312. Proceeding in this manner, we discover  $S_0 = \{0\}$ ,  $S_1 = \{1\}$ ,  $S_2 = \{2, 3\}$ ,  $S_3 = \{4, 5, 6\}$ , and  $S_4 = \{7\}$ .  $\Box$ 

Suppose that we have computed the  $S_k$ 's for position j-1 of the second string and we wish to modify them to apply to position j. We consider the set PLACES $(b_j)$ . For each r in PLACES $(b_j)$ , we consider whether we can improve some of the LCS's by adding the match between  $a_r$  and  $b_j$  to the LCS of  $a_1 \cdots a_{r-1}$  and  $b_1 \cdots b_j$ . That is, if both r-1 and r are in  $S_k$ , then all  $s \ge r$  in  $S_k$  really belong in  $S_{k+1}$  when  $b_j$  is considered. To see this we observe that we can obtain k matches between  $a_1 \cdots a_{r-1}$  and  $b_1 \cdots b_{j-1}$ , to which we add a match between  $a_r$  and  $b_j$ . We can modify  $S_k$  and  $S_{k+1}$  by the following steps.

- 1. FIND(r) to get  $S_k$ .
- 2. If FIND(r-1) is not  $S_k$ , then no benefit can be had by matching  $b_j$  with  $a_r$ . Skip the remaining steps and do not modify  $S_k$  or  $S_{k+1}$ .
- 3. If FIND $(r-1) = S_k$ , apply SPLIT $(S_k, S_k, S'_k, r)$  to separate from  $S_k$  those members greater than or equal to r.
- 4. MERGE( $S_k$ ,  $S_{k+1}$ ,  $S_{k+1}$ ) to move these elements into  $S_{k+1}$ .

It is important to consider the members of PLACES $(b_j)$  largest first. To see why, suppose for example that 7 and 9 are in PLACES $(b_j)$ , and before  $b_j$  is considered,  $S_3 = \{6, 7, 8, 9\}$  and  $S_4 = \{10, 11\}$ .

If we consider 7 before 9, we split  $S_3$  into  $S_3 = \{6\}$  and  $S'_3 = \{7, 8, 9\}$ , then make  $S_4 = \{7, 8, 9, 10, 11\}$ . If we then consider 9, we split  $S_4$  into  $S_4=\{7, 8\}$  and  $S'_4 = \{9, 10, 11\}$ , then merge 9, 10 and 11 into  $S_5$ . We have thus moved 9 from  $S_3$  to  $S_5$  by considering only one more position in the second string, representing an impossibility. Intuitively, what has happened is that we have erroneously matched  $b_i$  against both  $a_7$  and  $a_9$  in creating an imaginary LCS of length 5.

In Fig. 5.29, we see a sketch of the algorithm that maintains the sets  $S_k$  as we scan the second string. To determine the length of an LCS, we need only execute FIND(n) at the end.

	procedure LCS;	
	begin	
(1)	initialize $S_0 = \{0, 1,, n\}$ and $S_i = \emptyset$ for $i = 1, 2,$	.,n;
(2)	for $j := 1$ to n do { compute $S_k$ 's for position $j$ }	
(3)	for r in PLACES $(b_i)$ , largest first do begin	
(4)	k := FIND(r);	
(5)	if $k = FIND(r-1)$ then begin { r is not smallest i	$n S_k$
(6)	SPLIT( $S_k$ , $S_k$ , $S'_k$ , $r$ );	
(7)	MERGE $(S'_{k}, S_{k+1}, S_{k+1})$	
	end	
	end	

end; {LCS}

Fig. 5.29. Sketch of longest common subsequence program.

#### Time Analysis of the LCS Algorithm

As we mentioned earlier, the algorithm of Fig. 5.29 is a useful approach only if there are not too many matches between symbols of the two strings. The measure of the number of matches is

$$p = \sum_{j=1}^{m} | PLACES(b_j) |$$

where  $|PLACES(b_j)|$  denotes the number of elements in set  $PLACES(b_j)$ . In other words, p is the sum over all  $b_j$  of the number of positions in the first string that match  $b_j$ . Recall that in our discussion of file comparison, we expect p to be of the same order as m and n, the lengths of the two strings (files).

It turns out that the 2-3 tree is a good structure for the sets  $S_k$ . We can initialize these sets, as in line (1) of Fig. 5.29, in O(n) steps. The FIND operation requires an array to serve as a mapping from positions r to the leaf for r and also requires pointers to parents in the 2-3 tree. The name of the set, i.e., k for  $S_k$ , can be kept at the root, so we can execute FIND in  $O(\log n)$  steps by following parent pointers until we reach the root. Thus all executions of lines (4) and (5) together take  $O(p \log n)$  time, since those lines are each executed exactly once for each match found.

The MERGE operation of line (5) has the special property that every member of  $S'_k$  is lower than every member of  $S_{k+1}$ , and we can take advantage of this fact when using 2-3 trees for an implementation.<sup>†</sup> To begin the

<sup>†</sup> Strictly speaking we should use a different name for the MERGE operation, as the implementation we propose will not work to compute the arbitrary union of disjoint sets,

#### 5.6 AN ADT WITH MERGE AND SPLIT

MERGE, place the 2-3 tree for  $S'_k$  to the left of that for  $S_{k+1}$ . If both are of the same height, create a new root with the roots of the two trees as children. If  $S'_k$  is shorter, insert the root of that tree as the leftmost child of the leftmost node of  $S_{k+1}$  at the appropriate level. If this node now has four children, we modify the tree exactly as in the INSERT procedure of Fig. 5.20. An example is shown in Fig. 5.30. Similarly, if  $S_{k+1}$  is shorter, make its root the rightmost child of the rightmost node of  $S'_k$  at the appropriate level.



Fig. 5.30. Example of MERGE.

The SPLIT operation at r requires that we travel up the tree from leaf r, duplicating every interior node along the path and giving one copy to each of the two resulting trees. Nodes with no children are eliminated, and nodes with one child are removed and have that child inserted into the proper tree at the proper level.

**Example 5.9.** Suppose we split the tree of Fig. 5.30(b) at node 9. The two trees, with duplicated nodes, are shown in Fig. 5.31(a). On the left, the parent of 8 has only one child, so 8 becomes a child of the parent of 6 and 7. This parent now has three children, so all is as it should be; if it had four children, a new node would have been created and inserted into the tree. We need only eliminate nodes with zero children (the old parent of 8) and the chain of nodes with one child leading to the root. The parent of 6, 7, and 8 becomes the new root, as shown in Fig. 5.31(b). Similarly, in the right tree, 9 becomes a sibling of 10 and 11, and unnecessary nodes are eliminated, as is also shown in Fig. 5.31(b).  $\Box$ 

while keeping the elements sorted so operations like SPLIT and FIND can be performed.

ADVANCED SET REPRESENTATION METHODS



(b) Result of repairs.

Fig. 5.31. An example of SPLIT.

If we do the splitting and reorganization of the 2-3 tree bottom up, it can be shown by consideration of a large number of cases that  $O(\log n)$  steps suffices. Thus, the total time spent in lines (6) and (7) of Fig. 5.29 is  $O(p\log n)$ , and hence the entire algorithm takes  $O(p\log n)$  steps. We must add in the preprocessing time needed to compute and sort PLACES(a) for symbols a. As we mentioned, if the symbols a are "large" objects, this time can be much greater than any other part of the algorithm. As we shall see in Chapter 8, if the symbols can be manipulated and compared in single "steps," then  $O(n\log n)$  time suffices to sort the first string  $a_1a_2 \cdots a_n$  (actually, to sort objects  $(i, a_i)$  on the second field), whereupon PLACES(a) can be read off from this list in O(n) time. Thus, the length of the LCS can be computed in  $O(\max(n, p) \log n)$  time which, since  $p \ge n$  is normal, can be taken as  $O(p\log n)$ .

### EXERCISES

# Exercises

- 5.1 Draw all possible binary search trees containing the four elements 1, 2, 3, 4.
- 5.2 Insert the integers 7, 2, 9, 0, 5, 6, 8, 1 into a binary search tree by repeated application of the procedure INSERT of Fig. 5.3.
- 5.3 Show the result of deleting 7, then 2 from the final tree of Exercise 5.2.
- **\*5.4** When deleting two elements from a binary search tree using the procedure of Fig. 5.5, does the final tree ever depend on the order in which you delete them?
- 5.5 We with to keep track of all 5-character substrings that occur in a given string, using a trie. Show the trie that results when we insert the 14 substrings of length five of the string ABCDABACDEBA-CADEBA.
- \*5.6 To implement Exercise 5.5, we could keep a pointer at each leaf, which, say, represents string abcde, to the interior node representing the suffix bcde. That way, if the next symbol, say f, is received, we don't have to insert all of bcdef, starting at the root. Furthermore, having seen abcde, we may as well create nodes for bcde, cde, de, and e, since we shall, unless the sequence ends abruptly, need those nodes eventually. Modify the trie data structure to maintain such pointers, and modify the trie insertion algorithm to take advantage of this data structure.
- 5.7 Show the 2-3 tree that results if we insert into an empty set, represented as a 2-3 tree, the elements 5, 2, 7, 0, 3, 4, 6, 1, 8, 9.
- 5.8 Show the result of deleting 3 from the 2-3 tree that results from Exercise 5.7.
- 5.9 Show the successive values of the various  $S_i$ 's when implementing the LCS algorithm of Fig. 5.29 with first string *abacabada*, and second string *bdbacbad*.
- 5.10 Suppose we use 2-3 trees to implement the MERGE and SPLIT operations as in Section 5.6.
  - a) Show the result of splitting the tree of Exercise 5.7 at 6.
  - b) Merge the tree of Exercise 5.7 with the tree consisting of leaves for elements 10 and 11.
- 5.11 Some of the structures discussed in this chapter can be modified easily to support the MAPPING ADT. Write procedures MAKENULL, ASSIGN, and COMPUTE to operate on the following data structures.
  - a) Binary search trees. The "<" ordering applies to domain elements.

- b) 2-3 trees. At interior nodes, place only the key field of domain elements.
- 5.12 Show that in any subtree of a binary search tree, the minimum element is at a node without a left child.
- 5.13 Use Exercise 5.12 to produce a nonrecursive version of DELETE-MIN.
- 5.14 Write procedures ASSIGN, VALUEOF, MAKENULL and GETNEW for trie nodes represented as lists of cells.
- \*5.15 How do the trie (list of cells implementation), the open hash table, and the binary search tree compare for speed and for space utilization when elements are strings of up to ten characters?
- **\*5.16** If elements of a set are ordered by a "<" relation, then we can keep one or two elements (not just their keys) at interior nodes of a 2-3 tree, and we then do not have to keep these elements at the leaves. Write INSERT and DELETE procedures for 2-3 trees of this type.
- 5.17 Another modification we could make to 2-3 trees is to keep only keys at interior nodes, but do not require that the keys  $k_1$  and  $k_2$  at a node truly be the minimum keys of the second and third subtrees, just that all keys k of the third subtree satisfy  $k \ge k_2$ , all keys k of the second satisfy  $k_1 \le k < k_2$ , and all keys k of the first satisfy  $k < k_1$ .
  - a) How does this convention simplify the DELETE operation?
  - b) Which of the dictionary and mapping operations are made more complicated or less efficient?
- \*5.18 Another data structure that supports dictionaries with the MIN operation is the AVL tree (named for the inventors' initials) or height-balanced tree. These trees are binary search trees in which the heights of two siblings are not permitted to differ by more than one. Write procedures to implement INSERT and DELETE, while maintaining the AVL-tree property.
- 5.19 Write the Pascal program for procedure delete1 of Fig. 5.21.
- \*5.20 A finite automaton consists of a set of states, which we shall take to be the integers 1..n and a table transitions[state, input] giving a next state for each state and each input character. For our purposes, we shall assume that the input is always either 0 or 1. Further, certain of the states are designated accepting states. For our purposes, we shall assume that all and only the even numbered states are accepting. Two states p and q are equivalent if either they are the same state, or (i) they are both accepting or both nonaccepting, (ii) on input 0 they transfer to equivalent states, and (iii) on input 1 they transfer to equivalent states. Intuitively, equivalent states behave the same on all sequences of inputs; either both or neither lead to accepting states. Write a program using the MFSET operations that computes the sets of equivalent states of a given finite automaton.

**\*\*5.21** In the tree implementation of MFSET:

- a) Show that  $\Omega(n \log n)$  time is needed for certain lists of n operations if path compression is used but larger trees are permitted to be merged into smaller ones.
- b) Show that  $O(n\alpha(n))$  is the worst case running time for *n* operations if path compression is used, and the smaller tree is always merged into the larger.
- 5.22 Select a data structure and write a program to compute PLACES (defined in Section 5.6) in average time O(n) for strings of length n.
- **\*5.23** Modify the LCS procedure of Fig. 5.29 to compute the LCS, not just its length.
- \*5.24 Write a detailed SPLIT procedure to work on 2-3 trees.
- **\*5.25** If elements of a set represented by a 2-3 tree consist only of a key field, an element whose key appears at an interior node need not appear at a leaf. Rewrite the dictionary operations to take advantage of this fact and avoid storing any element at two different nodes.

#### **Bibliographic Notes**

Tries were first proposed by Fredkin [1960]. Bayer and McCreight [1972] introduced B-trees, which, as we shall see in Chapter 11, are a generalization of 2-3 trees. The first uses of 2-3 trees were by J. E. Hopcroft in 1970 (unpublished) for insertion, deletion, concatenation, and splitting, and by Ullman [1974] for a code optimization problem.

The tree structure of Section 5.5, using path compression and merging smaller into larger, was first used by M. D. Mcllroy and R. Morris to construct minimum-cost spanning trees. The performance of the tree implementation of MFSET's was analyzed by Fischer [1972] and by Hopcroft and Ullman [1973]. Exercise 5.21(b) is from Tarjan [1975].

The solution to the LCS problem of Section 5.6 is from Hunt and Szymanski [1977]. An efficient data structure for FIND, SPLIT, and the restricted MERGE (where all elements of one set are less than those of the other) is described in van Emde Boas, Kaas, and Zijlstra [1977].

Exercise 5.6 is based on an efficient algorithm for matching patterns developed by Weiner [1973]. The 2-3 tree variant of Exercise 5.16 is discussed in detail in Wirth [1976]. The AVL tree structure in Exercise 5.18 is from Adel'son-Vel'skii and Landis [1962].

# CHAPTER 10

# Algorithm Design Techniques

Over the years computer scientists have identified a number of general techniques that often yield effective algorithms in solving large classes of problems. This chapter presents some of the more important techniques, such as divide-and-conquer, dynamic programming, greedy techniques, backtracking, and local search. In trying to devise an algorithm to solve a give problem, it is often useful to ask a question such as "What kind of solution does divideand-conquer, dynamic programming, a greedy approach, or some other standard technique yield?"

It should be emphasized, however, that there are problems, such as the NP-complete problems, for which these or any other known techniques will not produce efficient solutions. When such a problem is encountered, it is often useful to determine if the inputs to the problem have special characteristics that could be exploited in trying to devise a solution, or if an easily found approximate solution could be used in place of the difficult-to-compute exact solution.

#### 10.1 Divide-and-Conquer Algorithms

Perhaps the most important, and most widely applicable, technique for designing efficient algorithms is a strategy called "divide-and-conquer." It consists of breaking a problem of size n into smaller problems in such a way that from solutions to the smaller problems we can easily construct a solution to the entire problem. We have already seen a number of applications of this technique, such as *mergesort* or binary search trees.

To illustrate the method consider the familiar "towers of Hanoi" puzzle. It consists of three pegs A, B, and C. Initially peg A has on it some number of disks, starting with the largest one on the bottom and successively smaller ones on top, as shown in Fig. 10.1. The object of the puzzle is to move the disks one at a time from peg to peg, never placing a larger disk on top of a smaller one, eventually ending with all disks on peg B.

One soon learns that the puzzle can be solved by the following simple algorithm. Imagine the pegs arranged in a triangle. On odd-numbered moves, move the smallest disk one peg clockwise. On even-numbered moves



Fig. 10.1. Initial position in towers of Hanoi puzzle.

make the only legal move not involving the smallest disk.

The above algorithm is concise, and correct, but it is hard to understand why it works, and hard to invent on the spur of the moment. Consider instead the following divide-and-conquer approach. The problem of moving the n smallest disks from A to B can be thought of as consisting of two subproblems of size n-1. First move the n-1 smallest disks from peg A to peg C, exposing the  $n^{th}$  smallest disk on peg A. Move that disk from A to B. Then move the n-1 smallest disks from C to B. Moving the n-1 smallest disks is accomplished by a recursive application of the method. As the n disks involved in the moves are smaller than any other disks, we need not concern ourselves with what is below them on pegs A, B, or C. Although the actual movement of individual disks is not obvious, and hand simulation is hard because of the stacking of recursive calls, the algorithm is conceptually simple to understand, to prove correct and, we would like to think, to invent in the first place. It is probably the ease of discovery of divide-and-conquer algorithms that makes the technique so important, although in many cases the algorithms are also more efficient than more conventional ones.<sup>†</sup>

# The Problem of Multiplying Long Integers

Consider the problem of multiplying two *n*-bit integers X and Y. Recall that the algorithm for multiplication of *n*-bit (or *n*-digit) integers usually taught in elementary school involves computing *n* partial products of size *n* and thus is an  $O(n^2)$  algorithm, if we count single bit or digit multiplications and additions as one step. One divide-and-conquer approach to integer multiplication would break each of X and Y into two integers of n/2 bits each as shown in

<sup>†</sup> In the towers of Hanoi case, the divide-and-conquer algorithm is really the same as the one given initially.

Fig. 10.2. (For simplicity we assume n is a power of 2 here.)



Fig. 10.2. Breaking *n*-bit integers into  $\frac{n}{2}$ -bit pieces.

The product of X and Y can now be written

$$XY = AC2^{n} + (AD + BC) 2^{n/2} + BD$$
(10.1)

If we evaluate XY in this straightforward way, we have to perform four multiplications of (n/2)-bit integers (AC, AD, BC, and BD), three additions of integers with at most 2n bits (corresponding to the three + signs in (10.1)), and two shifts (multiplication by  $2^n$  and  $2^{n/2}$ ). As these additions and shifts take O(n) steps, we can write the following recurrence for T(n), the total number of bit operations needed to multiply *n*-bit integers according to (10.1).

$$T(1) = 1$$
  

$$T(n) = 4T(n/2) + cn$$
(10.2)

Using reasoning like that in Example 9.4, we can take the constant c in (10.2) to be 1, so the driving function d(n) is just n, and then deduce that the homogeneous and particular solutions are both  $O(n^2)$ .

In the case that formula (10.1) is used to multiply integers, the asymptotic efficiency is thus no greater than for the elementary school method. But recall that for equations like (10.2) we get an asymptotic improvement if we decrease the number of subproblems. It may be a surprise that we can do so, but consider the following formula for multiplying X by Y.

$$XY = AC2^{n} + [(A-B)(D-C) + AC + BD]2^{n/2} + BD$$
(10.3)

Although (10.3) looks more complicated than (10.1) it requires only three multiplications of (n/2)-bit integers, AC, BD, and (A-B)(D-C), six additions or subtractions, and two shifts. Since all but the multiplications take O(n) steps, the time T(n) to multiply *n*-bit integers by (10.3) is given by

T(1) = 1

$$T(n) = 3T(n/2) + cn$$

whose solution is  $T(n) = O(n^{\log_2 3}) = O(n^{1.59})$ .

The complete algorithm, including the details implied by the fact that (10.3) requires multiplication of negative, as well as positive, (n/2)-bit

integers, is given in Fig. 10.3. Note that lines (8)-(11) are performed by copying bits, and the multiplication by  $2^n$  and  $2^{n/2}$  in line (16) by shifting. Also, the multiplication by s in line (16) simply introduces the proper sign into the result.

function mult (	X, Y, n: integer ): integer;
$\{X \text{ and } Y \text{ a}\}$	re signed integers $\leq 2^n$ .
n is a pow	er of 2. The function returns XY }
var	
s: integ	er; { holds the sign of XY }
m1, m2	, m3: integer; { hold the three products }
A, B, C	, D: integer; { hold left and right halves of X and Y }
begin	
(1)   s := sig	gn(X) * sign(Y);
(2)    X := al	$\operatorname{os}(X);$
(3) $Y := at$	os(Y); { make X and Y positive }
(4) <b>if</b> $n =$	l then
(5) if (.	X = 1) and $(Y = 1)$ then
(6)	return (s)
else	
(7)	return (0)
else beg	, in
(8) A:	= left $n/2$ bits of X;
(9) B:	= right $n/2$ bits of X;
(10) C:	= left $n/2$ bits of Y;
(11) D:	= right $n/2$ bits of Y;
(13) <i>m</i> 1	:= mult(A, C, n/2);
(14) m2	:= mult(A-B, D-C, n/2);
(15) m3	:= mult(B, D, n/2);
(16) ret	<b>IFT</b> $(s * (m1*2^{n} + (m1 + m2 + m3) * 2^{n/2} + m3))$
end	
end; { mult	}

Fig. 10.3. Divide-and-conquer integer multiplication algorithm.

Observe that the divide-and-conquer algorithm of Fig. 10.3 is asymptotically faster than the method taught in elementary school, taking  $O(n^{1.59})$  steps against  $O(n^2)$ . We may thus raise the question: if this algorithm is so superior why don't we teach it in elementary school? There are two answers. First, while easy to implement on a computer, the description of the algorithm is sufficiently complex that if we attempted to teach it in elementary school students would not learn to multiply. Furthermore, we have ignored constants of proportionality. While procedure *mult* of Fig. 10.3 is asymptotically superior to the usual method, the constants are such that for small problems (actually up to about 500 bits) the elementary school method is superior, and we rarely

ask elementary school children to multiply such numbers.

## **Constructing Tennis Tournaments**

The technique of divide-and-conquer has widespread applicability, not only in algorithm design but in designing circuits, constructing mathematical proofs and in other walks of life. We give one example as an illustration. Consider the design of a round robin tennis tournament schedule, for  $n = 2^k$  players. Each player must play every other player, and each player must play one match per day for n-1 days, the minimum number of days needed to complete the tournament.

The tournament schedule is thus an *n* row by n-1 column table whose entry in row *i* and column *j* is the player *i* must contend with on the *j*<sup>th</sup> day.

The divide-and-conquer approach constructs a schedule for one-half of the players. This schedule is designed by a recursive application of the algorithm by finding a schedule for one half of these players and so on. When we get down to two players, we have the base case and we simply pair them up.

Suppose there are eight players. The schedule for players 1 through 4 fills the upper left corner (4 rows by 3 columns) of the schedule being constructed. The lower left corner (4 rows by 3 columns) of the schedule must pit the high numbered players (5 through 8) against one another. This subschedule is obtained by adding 4 to each entry in the upper left.

We have now simplified the problem. All that remains is to have lowernumbered players play high-numbered players. This is easily accomplished by having players 1 through 4 play 5 through 8 respectively on day 4 and cyclically permuting 5 through 8 on subsequent days. The process is illustrated in Fig. 10.4. The reader should now be able to generalize the ideas of this figure to provide a schedule for  $2^k$  players for any k.

#### **Balancing Subproblems**

In designing algorithms one is always faced with various trade-offs. One rule that has emerged is that it is generally advantageous to balance competing costs wherever possible. For example in Chapter 5 we saw that the 2-3 tree balanced the costs of searching with those of inserting, while more straightforward methods take O(n) steps either for each lookup or for each insertion, even though the other operation can be done in a constant number of steps.

Similarly, for divide-and-conquer algorithms, we are generally better off if the subproblems are of approximately equal size. For example, insertion sort can be viewed as partitioning a problem into two subproblems, one of size 1 and one of size n-1, with a maximum cost of n steps to merge. This gives a recurrence

# T(n) = T(1) + T(n-1) + n

which has an  $O(n^2)$  solution. Mergesort, on the other hand, partitions the problems into two subproblems each of size n/2 and has  $O(n\log n)$  performance. As a general principle, we often find that partitioning a problem into



Fig. 10.4. A round-robin tournament for eight players.

equal or nearly equal subproblems is a crucial factor in obtaining good performance.

# 10.2 Dynamic Programming

Often there is no way to divide a problem into a small number of subproblems whose solution can be combined to solve the original problem. In such cases we may attempt to divide the problem into as many subproblems as necessary, divide each subproblem into smaller subproblems and so on. If this is all we do, we shall likely wind up with an exponential-time algorithm.

Frequently, however, there are only a polynomial number of subproblems, and thus we must be solving some subproblem many times. If instead we keep track of the solution to each subproblem solved, and simply look up the answer when needed, we would obtain a polynomial-time algorithm.

It is sometimes simpler from an implementation point of view to create a table of the solutions to all the subproblems we might ever have to solve. We fill in the table without regard to whether or not a particular subproblem is actually needed in the overall solution. The filling-in of a table of subproblems to get a solution to a given problem has been termed *dynamic programming*, a name that comes from control theory.

The form of a dynamic programming algorithm may vary, but there is the common theme of a table to fill and an order in which the entries are to be filled. We shall illustrate the techniques by two examples, calculating odds on a match like the World Series, and the "triangulation problem."

#### World Series Odds

Suppose two teams, A and B, are playing a match to see who is the first to win n games for some particular n. The World Series is such a match, with n = 4. We may suppose that A and B are equally competent, so each has a 50% chance of winning any particular game. Let P(i, j) be the probability that if A needs i games to win, and B needs j games, that A will eventually win the match. For example, in the World Series, if the Dodgers have won two games and the Yankees one, then i = 2, j = 3, and P(2, 3), we shall discover, is 11/16.

To compute P(i, j), we can use a recurrence equation in two variables. First, if i = 0 and j > 0, then team A has won the match already, so P(0, j) = 1. Similarly, P(i, 0) = 0 for i > 0. If i and j are both greater than 0, at least one more game must be played, and the two teams each win half the time. Thus, P(i, j) must be the average of P(i-1, j) and P(i, j-1), the first of these being the probability A will win the match if it wins the next game and the second being the probability A wins the match even though it loses the next game. To summarize:

$$P(i, j) = 1 \text{ if } i = 0 \text{ and } j > 0$$
  
= 0 if i > 0 and j = 0  
= (P(i-1, j) + P(i, j-1))/2 if i > 0 and j > 0 (10.4)

If we use (10.4) recursively as a function, we can show that P(i, j) takes no more than time  $O(2^{i+j})$ . Let T(n) be the maximum time taken by a call to P(i, j), where i+j = n. Then from (10.4),

T(1) = c

T(n) = 2T(n-1) + d

for some constants c and d. The reader may check by the means discussed in the previous chapter that  $T(n) \leq 2^{n-1}c + (2^{n-1}-1)d$ , which is  $O(2^n)$  or  $O(2^{i+j})$ .

We have thus proven an exponential upper bound on the time taken by the recursive computation of P(i, j). However, to convince ourselves that the recursive formula for P(i, j) is a bad way to compute it, we need to get a big-omega lower bound. We leave it as an exercise to show that when we call P(i, j), the total number of calls to P that gets made is  $\binom{i+j}{i}$ , the number of ways to choose *i* things out of i+j. If i = j, that number is  $\Omega(2^n/\sqrt{n})$ , where n = i+j. Thus, T(n) is  $\Omega(2^n/\sqrt{n})$ , and in fact, we can show it is  $O(2^n/\sqrt{n})$ also. While  $2^n/\sqrt{n}$  grows asymptotically more slowly than  $2^n$ , the difference is not great, and T(n) grows far too fast for the recursive calculation of P(i, j)to be practical.

The problem with the recursive calculation is that we wind up computing the same P(i, j) repeatedly. For example, if we want to compute P(2, 3), we compute, by (10.4), P(1, 3) and P(2, 2). P(1, 3) and P(2, 2) both require

the computation of P(1, 2), so we compute that value twice.

A better way to compute P(i, j) is to fill in the table suggested by Fig. 10.5. The bottom row is all 0's and the rightmost column all 1's by the first two lines of (10.4). By the last line of (10.4), each other entry is the average of the entry below it and the entry to the right. Thus, an appropriate way to fill in the table is to proceed in diagonals beginning at the lower right corner, and proceeding up and to the left along diagonals representing entries with a constant value of i+j, as suggested in Fig. 10.6. This program is given in Fig. 10.7, assuming it works on a two-dimensional array P of suitable size.







Fig. 10.6. Pattern of computation.

The analysis of function *odds* is easy. The loop of lines (4)-(5) takes O(s) time, and that dominates the O(1) time for lines (2)-(3). Thus, the outer loop takes time  $O(\sum_{s=1}^{n} s)$  or  $O(n^2)$ , where i+j = n. Thus dynamic pro-

	function odds (i, j: integer) : real;
	var
	s, k: integer;
	begin
(1)	for $s := 1$ to $i + j$ do begin
	{ compute diagonal of entries whose indices sum to s }
(2)	P[0, s] := 1.0;
(3)	P[s, 0] := 0.0;
(4)	for $k := 1$ to $s-1$ do
(5)	P[k, s-k] := (P[k-1, s-k] + P[k, s-k-1])/2.0
	end;
(6)	return $(P[i, j])$
	<b>end</b> ; { <i>odds</i> }

#### Fig. 10.7. Odds calculation.

gramming takes  $O(n^2)$  time, compared with  $O(2^n/\sqrt{n})$  for the straightforward approach. Since  $2^n/\sqrt{n}$  grows wildly faster than  $n^2$ , we would prefer dynamic programming to the recursive approach under essentially any circumstances.

#### **The Triangulation Problem**

As another example of dynamic programming, consider the problem of *tri*angulating a polygon. We are given the vertices of a polygon and a distance measure between each pair of vertices. The distance may be the ordinary (Euclidean) distance in the plane, or it may be an arbitrary cost function given by a table. The problem is to select a set of *chords* (lines between nonadjacent vertices) such that no two chords cross each other, and the entire polygon is divided into triangles. The total length (distance between endpoints) of the chords selected must be a minimum. We call such a set of chords a minimal triangulation.

**Example 10.1.** Figure 10.8 shows a seven-sided polygon and the (x, y) coordinates of its vertices. The distance function is the ordinary Euclidean distance. A triangulation, which happens not to be minimal, is shown by dashed lines. Its cost is the sum of the lengths of the chords  $(v_0, v_2), (v_0, v_3), (v_0, v_5)$ , and  $(v_3, v_5)$ , or  $\sqrt{8^2 + 16^2} + \sqrt{15^2 + 16^2} + \sqrt{22^2 + 2^2} + \sqrt{7^2 + 14^2} = 77.56$ .  $\square$ 

As well as being interesting in its own right, the triangulation problem has a number of useful applications. For example, Fuchs, Kedem, and Uselton [1977] used a generalization of the triangulation problem for the following purpose. Consider the problem of shading a two-dimensional picture of an object whose surface is defined by a collection of points in 3-space. The light source comes from a given direction, and the brightness of a point on the surface depends on the angles between the direction of light, the direction of the viewer's eye, and a perpendicular to the surface at that point. To estimate the direction of the surface at a point, we can compute a minimum triangulation



Fig. 10.8. A heptagon and a triangulation.

of the points defining the surface.

Each triangle defines a plane in a 3-space, and since a minimum triangulation was found, the triangles are expected to be very small. It is easy to find the direction of a perpendicular to a plane, so we can compute the light intensity for the points of each triangle, on the assumption that the surface can be treated as a triangular plane in a given region. If the triangles are not sufficiently small to make the light intensity look smooth, then local averaging can improve the picture.

Before proceeding with the dynamic programming solution to the triangulation problem, let us state two observations about triangulations that will help us design the algorithm. Throughout we assume we have a polygon with nvertices  $v_0, v_1, \ldots, v_{n-1}$ , in clockwise order.

Fact 1. In any triangulation of a polygon with more than three vertices, every pair of adjacent vertices is touched by at least one chord. To see this, suppose neither  $v_i$  nor  $v_{i+1}^{\dagger}$  were touched by a chord. Then the region that edge  $(v_i, v_{i+1})$  bounds would have to include edges  $(v_{i-1}, v_i)$ ,  $(v_{i+1}, v_{i+2})$  and at least one additional edge. This region then would not be a triangle.

Fact 2. If  $(v_i, v_j)$  is a chord in a triangulation, then there must be some  $v_k$ 

<sup>†</sup> In what follows, we take all subscripts to be computed modulo *n*. Thus, in Fig. 10.8,  $v_i$  and  $v_{i+1}$  could be  $v_6$  and  $v_0$ , respectively, since n = 7.

#### ALGORITHM DESIGN TECHNIQUES

such that  $(v_i, v_k)$  and  $(v_k, v_j)$  are each either edges of the polygon or chords. Otherwise,  $(v_i, v_j)$  would bound a region that was not a triangle.

To begin searching for a minimum triangulation, we pick two adjacent vertices, say  $v_0$  and  $v_1$ . By the two facts we know that in any triangulation, and therefore in the minimum triangulation, there must be a vertex  $v_k$  such that  $(v_1, v_k)$  and  $(v_k, v_0)$  are chords or edges in the triangulation. We must therefore consider how good a triangulation we can find after selecting each possible value for k. If the polygon has n vertices, there are a total of (n-2) choices to make.

Each choice of k leads to at most two subproblems, which we define to be polygons formed by one chord and the edges in the original polygon from one end of the chord to the other. For example, Fig. 10.9 shows the two subproblems that result if we select the vertex  $v_3$ .



Fig. 10.9. The two subproblems after selecting  $v_3$ .

Next, we must find minimum triangulations for the polygons of Fig. 10.9(a) and (b). Our first instinct is that we must again consider all chords emanating from two adjacent vertices. For example, in solving Fig. 10.9(b), we might consider choosing chord  $(v_3, v_5)$ , which leaves subproblem  $(v_0, v_3, v_5, v_6)$ , a polygon two of whose sides,  $(v_0, v_3)$  and  $(v_3, v_5)$ , are chords of the original polygon. This approach leads to an exponential-time algorithm.

However, by considering the triangle that involves the chord  $(v_0, v_k)$  we never have to consider polygons more than one of whose sides are chords of the original polygon. Fact 2 tells us that, in the minimal triangulation, the chord in the subproblem, such as  $(v_0, v_3)$  in Fig. 10.9(b), must make a triangle with one of the other vertices. For example, if we select  $v_4$ , we get the

triangle  $(v_0, v_3, v_4)$  and the subproblem  $(v_0, v_4, v_5, v_6)$  which has only one chord of the original polygon. If we try  $v_5$ , we get the subproblems  $(v_3, v_4, v_5)$  and  $(v_0, v_5, v_6)$ , with chords  $(v_3, v_5)$  and  $(v_0, v_5)$  only.

In general, define the subproblem of size s beginning at vertex  $v_i$ , denoted  $S_{is}$ , to be the minimal triangulation problem for the polygon formed by the s vertices beginning at  $v_i$  and proceeding clockwise, that is,  $v_i$ ,  $v_{i+1}$ , ...,  $v_{i+s-1}$ . The chord in  $S_{is}$  is  $(v_i, v_{i+s-1})$ . For example, Fig. 10.9(a) is  $S_{04}$  and Fig. 10.9(b) is  $S_{35}$ . To solve  $S_{is}$  we must consider the following three options.

- 1. We may pick vertex  $v_{i+s-2}$  to make a triangle with the chords  $(v_i, v_{i+s-1})$  and  $(v_i, v_{i+s-2})$  and third side  $(v_{i+s-2}, v_{i+s-1})$ , and then solve the subproblem  $S_{i,s-1}$ .
- 2. We may pick vertex  $v_{i+1}$  to make a triangle with the chords  $(v_i, v_{i+s-1})$  and  $(v_{i+1}, v_{i+s-1})$  and third side  $(v_i, v_{i+1})$ , and then solve the subproblem  $S_{i+1,s-1}$ .
- 3. For some k between 2 and s-3 we may pick vertex  $v_{i+k}$  and form a triangle with sides  $(v_i, v_{i+k})$ ,  $(v_{i+k}, v_{i+s-1})$ , and  $(v_i, v_{i+s-1})$  and then solve subproblems  $S_{i,k+1}$  and  $S_{i+k,s-k}$ .

If we remember that "solving" any subproblem of size three or less requires no action, we can summarize (1)-(3) by saying that we pick some k between 1 and s-2 and solve subproblems  $S_{i,k+1}$  and  $S_{i+k,s-k}$ . Figure 10.10 illustrates this division into subproblems.



Fig. 10.10. Division of  $S_{is}$  into subproblems.

If we use the obvious recursive algorithm implied by the above rules to solve subproblems of size four or more, then it is possible to show that each call on a subproblem of size s gives rise to a total of  $3^{s-4}$  recursive calls, if we "solve" subproblems of size three or less directly and count only calls on

#### ALGORITHM DESIGN TECHNIQUES

subproblems of size four or more. Thus the number of subproblems to be solved is exponential in s. Since our initial problem is of size n, where n is the number of vertices in the given polygon, the total number of steps performed by this recursive procedure is exponential in n.

Yet something is clearly wrong in this analysis, because we know that besides the original problem, there are only n(n-4) different subproblems that ever need to be solved. They are represented by  $S_{is}$ , where  $0 \le i < n$ and  $4 \le s < n$ . Evidently not all the subproblems solved by the recursive procedure are different. For example, if in Fig. 10.8 we choose chord  $(v_0, v_3)$ , and then in the subproblem of Fig. 10.9(b) we pick  $v_4$ , we have to solve subproblem  $S_{44}$ . But we would also have to solve this problem if we first picked chord  $(v_0, v_4)$ , or if we picked  $(v_1, v_4)$  and then, when solving subproblem  $S_{45}$ , picked vertex  $v_0$  to complete a triangle with  $v_1$  and  $v_4$ .

This suggests an efficient way to solve the triangulation problem. We make a table giving the cost  $C_{is}$  of triangulating  $S_{is}$  for all *i* and *s*. Since the solution to any given problem depends only on the solution to problems of smaller size, the logical order in which to fill in the table is in size order. That is, for sizes  $s = 4, 5, \ldots, n-1$  we fill in the minimum, cost for problems  $S_{is}$ , for all vertices *i*. It is convenient to include problems of size  $0 \le s < 4$  as well, but remember that  $S_{is}$  has cost 0 if s < 4.

By rules (1)-(3) above for finding subproblems, the formula for computing  $C_{is}$  for  $s \ge 4$  is:

$$C_{is} = \min_{1 \le k \le s-2} \left[ C_{i,k+1} + C_{i+k,s-k} + D(v_i, v_{i+k}) + D(v_{i+k}, v_{i+s-1}) \right] (10.5)$$

where  $D(v_p, v_q)$  is the length of the chord between vertices  $v_p$  and  $v_q$ , if  $v_p$ and  $v_q$  are not adjacent points on the polygon;  $D(v_p, v_q)$  is 0 if  $v_p$  and  $v_q$  are adjacent.

**Example 10.2.** Figure 10.11 holds the table of costs for  $S_{i,s}$  for  $0 \le i \le 6$  and  $4 \le s \le 6$ , based on the polygon and distances of Fig. 10.8. The costs for the rows with s < 3 are all zero. We have filled in the entry  $C_{07}$ , in column 0 and the row for s = 7. This entry, like all in that row, represents the triangulation of the entire polygon. To see that, just notice that we can, if we wish, consider the edge  $(v_0, v_6)$  to be a chord of a larger polygon and the polygon of Fig. 10.8 to be a subproblem of this polygon, which has a series of additional vertices extending clockwise from  $v_6$  to  $v_0$ . Note that the entire row for s = 7 has the same value as  $C_{07}$ , to within the accuracy of the computation.

Let us, as an example, show how the entry 38.09 in the column for i = 6 and row for s = 5 is filled in. According to (10.5) the value of this entry,  $C_{65}$ , is the minimum of three sums, corresponding to k = 1, 2, or 3. These sums are:

$$C_{62} + C_{04} + D(v_6, v_0) + D(v_0, v_3)$$
  

$$C_{63} + C_{13} + D(v_6, v_1) + D(v_1, v_3)$$
  

$$C_{64} + C_{22} + D(v_6, v_2) + D(v_2, v_3)$$

7	$C_{07} = 75.43$						
6	$C_{06} = 53.34$	$C_{16} = 55.22$	C <sub>26</sub> = 57.54	$C_{36} = 59.67$	$C_{46} = 59.78$	$C_{56} = 59.78$	$C_{66} = 63.61$
5	$C_{05} = 37.54$	$C_{15} = 31.81$	$C_{25} = 35.45$	$C_{35} = 37.74$	$C_{45} = 45.50$	$C_{55} = 39.98$	$C_{65} = 38.09$
4	$C_{04} = 16.16$	$C_{14} = 16.16$	$C_{24} = 15.65$	$C_{34} = 15.65$	$C_{44} = 22.09$	$C_{54} = 22.09$	$C_{64} = 17.89$
s	i = 0	1	2	3	4	5	6

**Fig. 10.11.** Table of  $C_{is}$ 's.

The distances we need are calculated from the coordinates of the vertices as:

$$D(v_2, v_3) = D(v_6, v_0) = 0$$

(since these are polygon edges, not chords, and are present "for free")

 $D(v_6, v_2) = 26.08$  $D(v_1, v_3) = 16.16$  $D(v_6, v_1) = 22.36$  $D(v_0, v_3) = 21.93$ 

The three sums above are 38.09, 38.52, and 43.97, respectively. We may conclude that the minimum cost of the subproblem  $S_{65}$  is 38.09. Moreover, since the first sum was smallest, we know that to achieve this minimum we must utilize the subproblems  $S_{62}$  and  $S_{04}$ , that is, select chord  $(\nu_0, \nu_3)$  and then solve  $S_{64}$  as best we can; chord  $(\nu_1, \nu_3)$  is the preferred choice for that subproblem.  $\Box$ 

There is a useful trick for filling out the table of Fig. 10.11 according to the formula (10.5). Each term of the min operation in (10.5) requires a pair of entries. The first pair, for k = 1, can be found in the table (a) at the "bottom" (the row for s = 2)† of the column of the element being computed, and (b) just below and to the right‡ of the element being computed. The second pair is (a) next to the bottom of the column, and (b) two positions down and to the right. Fig. 10.12 shows the two lines of entries we follow to get all the pairs of entries we need to consider simultaneously. The pattern — up the

<sup>&</sup>lt;sup>†</sup> Remember that the table of Fig. 10.11 has rows of 0's below those shown.

<sup>‡</sup> By "to the right" we mean in the sense of a table that wraps around. Thus, if we are at

the rightmost column, the column "to the right" is the leftmost column.

column and down the diagonal — is a common one in filling tables during dynamic programming.



Fig. 10.12. Pattern of table scan to compute one element.

#### Finding Solutions from the Table

While Fig. 10.11 gives us the cost of the minimum triangulation, it does not immediately give us the triangulation itself. What we need, for each entry, is the value of k that produced the minimum in (10.5). Then we can deduce that the solution consists of chords  $(v_i, v_{i+k})$ , and  $(v_{i+k}, v_{i+s-1})$  (unless one of them is not a chord, because k = 1 or k = s-2), plus whatever chords are implied by the solutions to  $S_{i,k+1}$  and  $S_{i+k,s-k}$ . It is useful, when we compute an element of the table, to include with it the value of k that gave the best solution.

**Example 10.3.** In Fig. 10.11, the entry  $C_{07}$ , which represents the solution to the entire problem of Fig. 10.8, comes from the terms for k = 5 in (10.5). That is, the problem  $S_{07}$  is split into  $S_{06}$  and  $S_{52}$ ; the former is the problem with six vertices  $v_0, v_1, \ldots, v_5$ , and the latter is a trivial "problem" of cost 0. Thus we introduce the chord  $(v_0, v_5)$  of cost 22.09 and must solve  $S_{06}$ .

The minimum cost for  $C_{06}$  comes from the terms for k = 2 in (10.5), whereby the problem  $S_{06}$  is split into  $S_{03}$  and  $S_{24}$ . The former is the triangle with vertices  $v_0$ ,  $v_1$ , and  $v_2$ , while the latter is the quadrilateral defined by  $v_2$ ,  $v_3$ ,  $v_4$ , and  $v_5$ .  $S_{03}$  need not be solved, but  $S_{24}$  must be, and we must include the costs of chords  $(v_0, v_2)$  and  $(v_2, v_5)$  which are 17.89 and 19.80, respectively. We find the minimum value for  $C_{24}$  is assumed when k = 1 in (10.5), giving us the subproblems  $C_{22}$  and  $C_{33}$ , both of which have size less than or equal to three and therefore cost 0. The chord  $(v_3, v_5)$  is introduced, with a cost of 15.65.  $\Box$ 



Fig. 10.13. A minimal cost triangulation.

# **10.3 Greedy Algorithms**

Consider the problem of making change. Assume coins of values  $25\phi$  (quarter),  $10\phi$  (dime),  $5\phi$  (nickel) and  $1\phi$  (penny), and suppose we want to return  $63\phi$  in change. Almost without thinking we convert this amount to two quarters, one dime and three pennies. Not only were we able to determine quickly a list of coins with the correct value, but we produced the shortest list of coins with that value.

The algorithm the reader probably used was to select the largest coin whose value was not greater than  $63\phi$  (a quarter), add it to the list and sub-tract its value from 63 getting  $38\phi$ . We then selected the largest coin whose value was not greater than  $38\phi$  (another quarter) and added it to the list, and so on.

This method of making change is a greedy algorithm. At any individual stage a greedy algorithm selects that option which is "locally optimal" in some particular sense. Note that the greedy algorithm for making change produces an overall optimal solution only because of special properties of the coins. If the coins had values 1¢, 5¢, and 11¢ and we were to make change of 15¢, the greedy algorithm would first select an 11¢ coin and then four 1¢ coins, for a total of five coins. However, three 5¢ coins would suffice.

We have already seen several greedy algorithms, such as Dijkstra's shortest path algorithm and Kruskal's minimum cost spanning tree algorithm. Dijkstra's shortest path algorithm is "greedy" in the sense that it always

#### ALGORITHM DESIGN TECHNIQUES

chooses the closest vertex to the source among those whose shortest path is not yet known. Kruskal's algorithm is also "greedy"; it picks from the remaining edges the shortest among those that do not create a cycle.

We should emphasize that not every greedy approach succeeds in producing the best result overall. Just as in life, a greedy strategy may produce a good result for a while, yet the overall result may be poor. As an example, we might consider what happens when we allow negative-weight edges in Dijkstra's and Kruskal's algorithms. It turns out that Kruskal's spanning tree algorithm is not affected; it still produces the minimum cost tree. But Dijkstra's algorithm fails to produce shortest paths in some cases.

**Example 10.4.** We see in Fig. 10.14 a graph with a negative cost edge between b and c. If we apply Dijkstra's algorithm with source s, we correctly discover first that the minimum path to a has length 1. Now, considering only edges from s or a to b or c, we expect that b has the shortest path from s, namely  $s \rightarrow a \rightarrow b$ , of length 3. We then discover that c has a shortest path from s of length 1.



Fig. 10.14. Graph with negative weight edge.

However, our "greedy" selection of b before c was wrong from a global point of view. It turns out that the path  $s \rightarrow a \rightarrow c \rightarrow b$  has length only 2, so our minimum distance of 3 for b was wrong.<sup>†</sup>

# **Greedy Algorithms as Heuristics**

For some problems no known greedy algorithm produces an optimal solution, yet there are greedy algorithms that can be relied upon to produce "good" solutions with high probability. Frequently, a suboptimal solution with a cost a few percent above optimal is quite satisfactory. In these cases, a greedy algorithm often provides the fastest way to get a "good" solution. In fact, if the problem is such that the only way to get an optimal solution is to use an

<sup>†</sup> In fact, we should be careful what we mean by "shortest path" when there are negative edges. If we allow negative cost cycles, then we could traverse such a cycle repeatedly to get arbitrarily large negative distances, so presumably we want to restrict ourselves to acyclic paths.

## 10.3 GREEDY ALGORITHMS

exhaustive search technique, then a greedy algorithm or other heuristic for getting a good, but not necessarily optimal, solution may be our only real choice.

**Example 10.5.** Let us introduce a famous problem where the only known algorithms that produce optimal solutions are of the "try-all-possibilities" variety and can have running times that are exponential in the size of the input. The problem, called the *traveling salesman problem*, or *TSP*, is to find, in an undirected graph with weights on the edges, a *tour* (a simple cycle that includes all the vertices) the sum of whose edge-weights is a minimum. A tour is often called a Hamilton (or Hamiltonian) cycle.

Figure 10.15(a) shows one instance of the traveling salesman problem, a graph with six vertices (often called "cities"). The coordinates of each vertex are given, and we take the weight of each edge to be its length. Note that, as is conventional with the TSP, we assume all edges exist, that is, the graph is complete. In more general instances, where the weight of edges is not based on Euclidean distance, we might find a weight of infinity on an edge that really was not present.

Figure 10.15(b)-(e) shows four tours of the six "cities" of Fig. 10.15(a). The reader might ponder which of these four, if any, is optimal. The lengths of these four tours are 50.00, 49.73, 48.39, and 49.78, respectively; (d) is the shortest of all possible tours.



Fig. 10.15. An instance of the traveling salesman problem.

The TSP has a number of practical applications. As its name implies, it

#### ALGORITHM DESIGN TECHNIQUES

can be used to route a person who must visit a number of points and return to his starting point. For example, the TSP has been used to route collectors of coins from pay phones. The vertices are the phones and the "home base." The cost of each edge is the travel time between the two points in question.

Another "application" of the TSP is in solving the *knight's tour problem*: find a sequence of moves whereby a knight can visit each square of the chessboard exactly once and return to its starting point. Let the vertices be the chessboard squares and let the edges between two squares that are a knight's move apart have weight 0; all other edges have weight infinity. An optimal tour has weight 0 and must be a knight's tour. Surprisingly, good heuristics for the TSP have no trouble finding knight's tours, although finding one "by hand" is a challenge.

The greedy algorithm for the TSP we have in mind is a variant of Kruskal's algorithm. Like that algorithm, we shall consider edges shortest first. In Kruskal's algorithm we accept an edge in its turn if it does not form a cycle with the edges already accepted, and we reject the edge otherwise. For the TSP, the acceptance criterion is that an edge under consideration, together with already selected edges,

- 1. does not cause a vertex to have degree three or more, and
- 2. does not form a cycle, unless the number of selected edges equals the number of vertices in the problem.

Collections of edges selected under these criteria will form a collection of unconnected paths, until the last step, when the single remaining path is closed to form a tour.

In Fig. 10.15(a), we would first pick edge (d, e), since it is the shortest, having length 3. Then we consider edges (b, c), (a, b), and (e, f), all of which have length 5. It doesn't matter in which order we consider them; all meet the conditions for selection, and we must select them if we are to follow the greedy approach. Next shortest edge is (a, c), with length 7.08. However, this edge would form a cycle with (a, b) and (b, c), so we reject it. Edge (d, f) is next rejected on similar grounds. Edge (b, e) is next to be considered, but it must be rejected because it would raise the degrees of b and e to three, and could then never form a tour with what we had selected. Similarly we reject (b, d). Next considered is (c, d), which is accepted. We now have one path,  $a \rightarrow b \rightarrow c \rightarrow d \rightarrow e \rightarrow f$ , and eventually accept (a, f) to complete the tour. The resulting tour is Fig. 10.15(b), which is fourth best of all the tours, but less than 4% more costly than the optimal.  $\Box$ 

# 10.4 Backtracking

Sometimes we are faced with the task of finding an optimal solution to a problem, yet there appears to be no applicable theory to help us find the optimum, except by resorting to exhaustive search. We shall devote this section to a systematic, exhaustive searching technique called backtracking and a technique called alpha-beta pruning, which frequently reduces the search substantially.

# 10.4 BACKTRACKING

Consider a game such as chess, checkers, or tic-tac-toe, where there are two players. The players alternate moves, and the state of the game can be represented by a board position. Let us assume that there are a finite number of board positions and that the game has some sort of stopping rule to ensure termination. With each such game, we can associate a tree called the *game tree*. Each node of the tree represents a board position. With the root we associate the starting position. If board position x is associated with node n, then the children of n correspond to the set of allowable moves from board position x, and with each child is associated the resulting board position. For example, Fig. 10.16 shows part of the tree for tic-tac-toe.



Fig. 10.16. Part of the tic-tac-toe game tree.

The leaves of the tree correspond to board positions where there is no move, either because one of the players has won or because all squares are filled and a draw resulted. We associate a value with each node of the tree.

First we assign values to the leaves. Say the game is tic-tac-toe. Then a leaf is assigned -1, 0 or 1 depending on whether the board position corresponds to a loss, draw or win for player 1 (playing X).

The values are propagated up the tree according to the following rule. If a node corresponds to a board position where it is player 1's move, then the value is the maximum of the values of the children of that node. That is, we assume player 1 will make the move most favorable to himself i.e., that which produces the highest-valued outcome. If the node corresponds to player 2's move, then the value is the minimum of the values of the children. That is, we assume player 2 will make his most favorable move, producing a loss for player 1 if possible, and a draw as next preference.

**Example 10.6.** The values of the boards have been marked in Fig. 10.16. The leaves that are wins for O get value -1, while those that are draws get 0, and wins for X get +1. Then we proceed up the tree. On level 8, where only one empty square remains, and it is X's move, the values for the unresolved boards is the "maximum" of the one child at level 9.

On level 7, where it is O's move and there are two choices, we take as a value for an interior node the minimum of the values of its children. The leftmost board shown on level 7 is a leaf and has value 1, because it is a win for X. The second board on level 7 has value  $\min(0, -1) = -1$ , while the third board has value  $\min(0, 1) = 0$ . The one board shown at level 6, it being X's move on that level, has value  $\max(1, -1, 0) = 1$ , meaning that there is some choice X can make that will win; in this case the win is immediate.  $\Box$ 

Note that if the root has value 1, then player 1 has a winning strategy. Whenever it is his turn he is guaranteed that he can select a move that leads to a board position of value 1. Whenever it is player 2's move he has no real choice but to select a moving leading to a board position of value 1, a loss for him. The fact that a game is assumed to terminate guarantees an eventual win for the first player. If the root has value 0, as it does in tic-tac-toe, then neither player has a winning strategy but can only guarantee himself a draw by playing as well as possible. If the root has value -1, then player 2 has a winning strategy.

#### **Payoff Functions**

The idea of a game tree, where nodes have values -1, 0, and 1, can be generalized to trees where leaves are given any number (called the *payoff*) as a value, and the same rules for evaluating interior nodes applies: take the maximum of the children on those levels where player 1 is to move, and the minimum of the children on levels where player 2 moves.

As an example where general payoffs are useful, consider a complex game, like chess, where the game tree, though finite, is so huge that evaluating it from the bottom up is not feasible. Chess programs work, in essence, by building for each board position from which it must move, the game tree with that board as root, extending downward for several levels; the exact

#### 10.4 BACKTRACKING

number of levels depends on the speed with which the computer can work. As most of the leaves of the tree will be ambiguous, neither wins, losses, nor draws, each program uses a function of board positions that attempts to estimate the probability of the computer winning in that position. For example, the difference in material figures heavily into such an estimation, as do such factors as the defensive strength around the kings. By using this payoff function, the computer can estimate the probability of a win after making each of its possible next moves, on the assumption of subsequent best play by each side, and chose the move with the highest payoff.<sup>†</sup>

# **Implementing Backtrack Search**

Suppose we are given the rules for a game,<sup>‡</sup> that is, its legal moves and rules for termination. We wish to construct its game tree and evaluate the root. We could construct the tree in the obvious way, and then visit the nodes in postorder. The postorder traversal assures that we visit an interior node n after all its children, whereupon we can evaluate n by taking the min or max of the values of its children, as appropriate.

The space to store the tree can be prohibitively large, but by being careful we need never store more than one path, from the root to some node, at any one time. In Fig. 10.17 we see the sketch of a recursive program that represents the path in the tree by the sequence of active procedure calls at any time. That program assumes the following:

- 1. Payoffs are real numbers in a limited range, for example -1 to +1.
- 2. The constant  $\infty$  is larger than any positive payoff and its negation is smaller than any negative payoff.
- 3. The type modetype is declared

#### type

# modetype = (MIN, MAX)

- 4. There is a type board type declared in some manner suitable for the representation of board positions.
- 5. There is a function *payoff* that computes the payoff for any board that is a *leaf* (i.e., won, lost, or drawn position).

† Incidentally, some of the other things good chessplaying programs do are:

- 1. Use heuristics to eliminate from consideration certain moves that are unlikely to be good. This helps expand the tree to more levels in a fixed time.
- 2. Expand "capture chains", which are sequences of capturing moves beyond the last level to which the tree is normally expanded. This helps estimate the relative material strength of positions more accurately.

3. Prune the tree search by alpha-beta pruning, as discussed later in this section.

<sup>‡</sup> We should not imply that only "games" can be solved in this manner. As we shall see in subsequent examples, the "game" could really represent the solution to a practical problem.

	function search (B: boardtype; mode: modetype) : real;
	{ evaluates the payoff for board B, assuming it is
	player 1's move if $mode = MAX$ and player 2's move
	if $mode = MIN$ . Returns the payoff }
	var
	C: boardtype; { a child of board $B$ }
	value: real; { temporary minimum or maximum value }
	begin
(1)	if B is a leaf then
(2)	return $(payoff(B))$
	else begin
	{ initialize minimum or maximum value of children }
(3)	if $mode = MAX$ then
(4)	value := $-\infty$
	else
(5)	value := $\infty$ ;
(6)	for each child $C$ of board $B$ do
(7)	if $mode = MAX$ then
(8)	value := $max(value, search(C, MIN))$
	else
(9)	value := $\min(value, search(C, MAX));$
10)	return (value)
	end
	end; { search }

Fig. 10.17. Recursive backtrack search program.

Another implementation we might consider is to use a nonrecursive program that keeps a stack of boards corresponding to the sequence of active calls to search. The techniques discussed in Section 2.6 can be used to construct such a program.

# **Alpha-Beta Pruning**

There is a simple observation that allows us to eliminate from consideration much of a typical game tree. In terms of Fig. 10.17, the for-loop of line (6) can skip over certain children, often many of the children. Suppose we have a node n, as in Fig. 10.18, and we have already determined that  $c_1$ , the first of n's children, has a value of 20. As n is a max node, we know its value is at least 20. Now suppose that continuing with our search we find that d, a child of  $c_2$  has value 15. As  $c_2$ , another child of n, is a min node, we know the value of  $c_2$  cannot exceed 15. Thus, whatever value  $c_2$  has, it cannot affect the value of n or any parent of n.

It is thus possible in the situation of Fig. 10.18, to skip consideration of the children of  $c_2$  that we have not already examined. The general rules for

(1



Fig. 10.18. Pruning the children of a node.

skipping or "pruning" nodes involves the notion of final and tentative values for nodes. The *final* value is what we have simply been calling the "value." A *tentative* value is an upper bound on the value of a min node, or a lower bound on the value of a max node. The rules for computing final and tentative values are the following.

- 1. If all the children of a node *n* have been considered or pruned, make the tentative value of *n* final.
- 2. If a max node *n* has tentative value  $v_1$  and a child with final value  $v_2$ , then set the tentative value of *n* to max $(v_1, v_2)$ . If *n* is a min node, set its tentative value to min $(v_1, v_2)$ .
- 3. If p is a min node with parent q (a max node), and p and q have tentative values  $v_1$  and  $v_2$ , respectively, with  $v_1 \le v_2$ , then we may prune all the unconsidered children of p. We may also prune the unconsidered children of p if p is a max node (and therefore q is a min node) and  $v_2 \le v_1$ .

**Example 10.7.** Consider the tree in Fig. 10.19. Assuming values for the leaves as shown, we wish to calculate the value for the root. We begin a post-order traversal. After reaching node D, by rule (2) we assign a tentative value of 2, which is the final value of D, to node C. We then search E and return to C and then to B. By rule (1), the final value of C is fixed at 2 and the value of B is tentatively set to 2. The search continues down to G and then back to F. The value F is tentatively set to 6. By rule (3), with p and q equal to F and B, respectively, we may prune H. That is, there is no need to search node H, since the tentative value of F can never decrease and it is already greater than the value of B, which can never increase.





Continuing our example, A is assigned a tentative value of 2 and the search proceeds to K. J is assigned a tentative value of 8. L does not determine the value of max node J. I is assigned a tentative value of 8. The search goes down to N, and M is assigned a tentative value of 5. Node O must be searched, since 5, the tentative value of M, is less than the tentative value of I. The tentative values of I and A are revised, and the search goes down to R. Eventually R and S are searched, and P is assigned a tentative value of 4. We need not search T or below, since that can only lower P's value and it is already too low to affect the value of A.  $\Box$ 

#### **Branch-and-Bound Search**

Games are not the only sorts of "problems" that can be solved exhaustively by searching a complete tree of possibilities. A wide variety of problems where we are asked to find a minimum or maximum configuration of some sort are amenable to solution by backtracking search over a tree of all possibilities. The nodes of the tree can be thought of as sets of configurations, and the children of a node n each represent a subset of the configurations that n represents. Finally, the leaves each represent single configurations, or solutions to the problem, and we may evaluate each such configuration to see if it is the best solution found so far.

If we are reasonably clever in how we design the search, the children of a node will each represent far fewer configurations than the node itself, so we need not go to too great a depth before reaching leaves. Lest this notion of searching appear too vague, let us take a concrete example.

Example 10.8. Recall from the previous section our discussion of the

#### 10.4 BACKTRACKING

traveling salesman problem. There we gave a "greedy algorithm" for finding a good but not necessarily optimum tour. Now let us consider how we might find the optimum tour by systematically considering all tours. One way is to consider all permutations of the nodes, and evaluate the tour that visits the nodes in that order, remembering the best found so far. The time for such an approach is O(n!) on an *n* node graph, since we must consider (n-1)! different permutations,<sup>†</sup> and each permutation takes O(n) time to evaluate.

We shall consider a somewhat different approach that is no better than the above in the worst case, but on the average, when coupled with a technique called "branch-and-bound" that we shall discuss shortly, produces the answer far more rapidly than the "try all permutations" method. Start constructing a tree, with a root that represents all tours. Tours are what we called "configurations" in the prefatory material. Each node has two children, and the tours that a node represents are divided by these children into two groups — those that have a particular edge and those that do not. For example, Fig. 10.20 shows portions of the tree for the TSP instance from Fig. 10.15(a).

In Fig. 10.20 we have chosen to consider the edges in lexicographic order  $(a, b), (a, c), \ldots, (a, f), (b, c), \ldots, (b, f), (c, d)$ , and so on. We could, of course pick any other order. Observe that not every node in the tree has two children. We can eliminate some children because the edges selected do not form part of a tour. Thus, there is no node for "tours containing (a, b), (a, c), and (a, d)," because a would have degree 3 and the result would not be a tour. Similarly, as we go down the tree we shall eliminate some nodes because some city would have degree less than 2. For example, we shall find no node for tours without (a, b), (a, c), (a, d), or (a, e).  $\Box$ 

# **Bounding Heuristics Needed for Branch-and-Bound**

Using ideas similar to those in alpha-beta pruning, we can eliminate far more nodes of the search tree than would be suggested by Example 10.8. Suppose, to be specific, that our problem is to minimize some function, e.g., the cost of a tour in the TSP. Suppose also that we have a method for getting a lower bound on the cost of any solution among those in the set of solutions represented by some node n. If the best solution found so far costs less than the lower bound for node n, we need not explore any of the nodes below n.

**Example 10.9.** We shall discuss one way to get a lower bound on certain sets of solutions for the TSP, those sets represented by nodes in a tree of solutions as suggested in Fig. 10.20. First of all, suppose we wish a lower bound on all solutions to a given instance of the TSP. Observe that the cost of any tour can be expressed as one half the sum over all nodes n of the cost of the two tour edges adjacent to n. This remark leads to the following rule. The sum

<sup>&</sup>lt;sup>†</sup> Note that we need not consider all n! permutations, since the starting point of a tour is immaterial. We may therefore consider only those permutations that begin with 1.

ALGORITHM DESIGN TECHNIQUES



Fig. 10.20. Beginning of a solution tree for a TSP instance.

of the two tour edges adjacent to node n is no less than the sum of the two edges of least cost adjacent to n. Thus, no tour can cost less than one half the sum over all nodes n of the two lowest cost edges incident upon n.

For example, consider the TSP instance in Fig. 10.21. Unlike the instance in Fig. 10.15, the distance measure for edges is not Euclidean; that is, it bears no relation to the distance in the plane between the cities it connects. Such a cost measure might be traveling time or fare, for example. In this instance, the least cost edges adjacent to node a are (a, d), and (a, b), with a total cost of 5. For node b, we have (a, b) and (b, e), with a total cost of 6. Similarly, the two lowest cost edges adjacent to c, d, and e, total 8, 7, and 9, respectively. Our lower bound on the cost of a tour is thus (5+6+8+7+9)/2 = 17.5.




Now suppose we want a lower bound on the cost of a subset of tours defined by some node in the search tree. If the search tree is constructed as in Example 10.8, each node represents tours defined by a set of edges that must be in the tour and a set of edges that may not be in the tour. These *constraints* alter our choices for the two lowest-cost edges at each node. Obviously an edge constrained to be in any tour must be included among the two edges selected, no matter whether they are or are not lowest or second lowest in cost.<sup>†</sup> Similarly, an edge constrained to be out cannot be selected, even if its cost is lowest.

Thus, if we are constrained to include edge (a, e), and exclude (b, c), the two edges for node a are (a, d) and (a, e), with a total cost of 9. For bwe select (a, b) and (b, e), as before, with a total cost of 6. For c, we cannot select (b, c), and so select (a, c) and (c, d), with a total cost of 9. For dwe select (a, d) and (c, d), as before, while for e we must select (a, e), and choose to select (b, e). The lower bound for these constraints is thus (9+6+9+7+10)/2 = 20.5.  $\Box$ 

Now let us construct the search tree along the lines suggested in Example 10.8. We consider the edges in lexicographic order, as in that example. Each

<sup>&</sup>lt;sup>†</sup> The rules for constructing the search tree will be seen to eliminate any set of constraints that cannot yield any tour, e.g., because three edges adjacent to one node are required to be in the tour.

time we *branch*, by considering the two children of a node, we try to infer additional decisions regarding which edges must be included or excluded from tours represented by those nodes. The rules we use for these inference are:

- 1. If excluding an edge (x, y) would make it impossible for x or y to have as many as two adjacent edges in the tour, then (x, y) must be included.
- 2. If including (x, y) would cause x or y to have more than two edges adjacent in the tour, or would complete a non-tour cycle with edges already included, then (x, y) must be excluded.

When we branch, after making what inferences we can, we compute lower bounds for both children. If the lower bound for a child is as high or higher than the lowest cost tour found so far, we can "prune" that child and need not construct or consider its descendants. Interestingly, there are situations where the lower bound for a node n is lower than the best solution so far, yet both children of n can be pruned because their lower bounds exceed the cost of the best solution so far.

If neither child can be pruned, we shall, as a heuristic, consider the child with the smaller lower bound first, in the hope of more rapidly reaching a solution that is cheaper than the one so far found best.<sup>†</sup> After considering one child, we must consider again whether its sibling can be pruned, since a new best solution may have been found. For the instance of Fig. 10.21, we get the search tree of Fig. 10.22. To interpret nodes of that tree, it helps to understand that the capital letters are names of the search tree nodes. The numbers are the lower bounds, and we list the constraints applying to that node but none of its ancestors by writing xy if edge (x, y) must be included and  $\overline{xy}$  if (x, y) must be excluded. Also note that the constraints introduced at a node apply to all its descendants. Thus to get all the constraints applying at a node we must follow the path from that node to the root.

Lastly, let us remark that as for backtrack search in general, we construct the tree one node at a time, retaining only one path, as in the recursive algorithm of Fig. 10.17, or its nonrecursive counterpart. The nonrecursive version is probably to be preferred, so that we can maintain the list of constraints conveniently on a stack.

**Example 10.10.** Figure 10.22 shows the search tree for the TSP instance of Fig. 10.21. To see how it is constructed, we begin at the root A of Fig. 10.22. The first edge in lexicographic order is (a, b), so we consider the two children B and C, corresponding to the constraints ab and ab, respectively. There is, as yet, no "best solution so far," so we shall consider both B and C eventually.<sup>‡</sup> Forcing (a, b) to be included does not raise the lower bound, but

<sup>‡</sup> We could start with some heuristically found solution, say the greedy one, although that would not affect this example. The greedy solution for Fig. 10.21 has cost 21.

<sup>&</sup>lt;sup>†</sup> An alternative is to use a heuristic to obtain a good solution using the constraints required for each child. For example, the reader should be able to modify the greedy TSP algorithm to respect constraints.

## 10.4 BACKTRACKING

excluding it raises the bound to 18.5, since the two cheapest legal edges for nodes a and b now total 6 and 7, respectively, compared with 5 and 6 with no constraints. Following our heuristic, we shall consider the descendants of node B first.

The next edge in lexicographic order is (a, c). We thus introduce children D and E corresponding to tours where (a, c) is included and excluded, respectively. In node D, we can infer that neither (a, d) nor (a, e) can be in a tour, else a would have too many edges incident. Following our heuristic we consider E before D, and branch on edge (a, d). The children F and G are introduced with lower bounds 18 and 23, respectively. For each of these children we know about three of the edges incident upon a, and so can infer something about the remaining edge (a, e).

Consider the children of F first. The first remaining edge in lexicographic order is (b, c). If we include (b, c), then, as we have included (a, b), we cannot include (b, d) or (b, e). As we have eliminated (a, e) and (b, e), we must have (c, e) and (d, e). We cannot have (c, d) or c and d would have three incident edges. We are left with one tour (a, b, c, e, d, a), whose cost is 23. Similarly, node I, where (b, c) is excluded, can be proved to represent only the tour (a, b, e, c, d, a), of cost 21. That tour has the lowest cost found so far.

We now backtrack to E and consider its second child, G. But G has a lower bound of 23, which exceeds the best cost so far, 21. Thus we prune G. We now backtrack to B and consider its other child, D. The lower bound on D is 20.5, but since costs are integers, we know no tour represented by D can have cost less than 21. Since we already have a tour that cheap, we need not explore the descendants of D, and therefore prune D. Now we backtrack to A and consider its second child, C.

At the level of node C, we have only considered edge (a, b). Nodes J and K are introduced as children of C. J corresponds to those tours that have (a, c) but not (a, b), and its lower bound in 18.5. K corresponds to tours having neither (a, b) nor (a, c), and we may infer that those tours have (a, d) and (a, e). The lower bound for K is 21, and we may immediately prune K, since we already know a tour that is low in cost.

We next consider the children of J, which are L and M, and we prune M because its lower bound exceeds the best tour cost so far. The children of L are N and P, corresponding to tours that have (b, c), and that exclude (b, c). By considering the degree of nodes b and c, and remembering that the selected edges cannot form a cycle of fewer than all five cities, we can infer that nodes N and P each represent single tours. One of these, (a, c, b, e, d, a), has the lowest cost of any tour, 19. We have explored or pruned the entire tree and therefore end.  $\Box$ 



# Fig. 10.22. Search tree for TSP solution.

## **10.5 Local Search Algorithms**

Sometimes the following strategy will produce an optimal solution to a problem.

- 1. Start with a random solution.
- 2. Apply to the current solution a transformation from some given set of transformations to improve the solution. The improvement becomes the new "current" solution.
- 3. Repeat until no transformation in the set improves the current solution.

The resulting solution may or may not be optimal. In principle, if the

## 10.5 LOCAL SEARCH ALGORITHMS

"given set of transformations" includes all the transformations that take one solution and replace it by any other, then we shall never stop until we reach an optimal solution. But then the time to apply (2) above is the same as the time needed to examine all solutions, and the whole approach is rather pointless.

The method makes sense when we can restrict our set of transformations to a small set, so we can consider all transformations in a short time; perhaps  $O(n^2)$  or  $O(n^3)$  transformations should be allowed when the problem is of "size" n. If the transformation set is small, it is natural to view the solutions that can be transformed to one another in one step as "close." The transformations are called "local transformations," and the method is called *local* search.

**Example 10.11.** One problem we can solve exactly by local search is the minimal spanning tree problem. The local transformations are those in which we take some edge not in the current spanning tree, add it to the tree, which must produce a unique cycle, and then eliminate exactly one edge of the cycle (presumably that of highest cost) to form a new tree.

For example, consider the graph of Fig. 10.21. We might start with the tree shown in Fig. 10.23(a). One transformation we could perform is to add edge (d, e) and remove another edge in the cycle formed, which is (e, a, c, d, e). If we remove edge (a, e), we decrease the cost of the tree from 20 to 19. That transformation can be made, leaving the tree of Fig. 10.23(b), to which we again try to apply an improving transformation. One such is to insert edge (a, d) and delete edge (c, d) from the cycle formed. The result is shown in Fig. 10.23(c). Then we might introduce (a, b) and delete (b, c) as in Fig. 10.23(d), and subsequently introduce (b, e) in favor of (d, e). The resulting tree of Fig. 10.23(e) is minimal. We can check that every edge not in that tree has the highest cost of any edge in the cycle it forms. Thus no transformation is applicable to Fig. 10.23(e).  $\Box$ 

The time taken by the algorithm of Example 10.11 on a graph of n nodes and e edges depends on the number of times we need to improve the solution. Just testing that no transformation is applicable could take O(ne) time, since eedges must be tried, and each could form a cycle of length nearly n. Thus the algorithm is not as good as Prim's or Kruskal's algorithms, but serves as an example where an optimal solution can be obtained by local search.

## Local Search Approximation Algorithms

Local search algorithms have had their greatest effectiveness as heuristics for the solution to problems whose exact solutions require exponential time. A common method of search is to start with a number of random solutions, and apply the local transformations to each, until reaching a *locally optimal* solution, one that no transformation can improve. We shall frequently reach different locally optimal solutions, from most or all of the random starting solutions, as suggested in Fig. 10.24. If we are lucky, one of them will be



Fig. 10.23. Local search for a minimal spanning tree.

globally optimal, that is, as good as any solution.

In practice, we may not find a globally optimal solution as suggested in Fig. 10.24, since the number of locally optimal solutions may be enormous. However, we may at least choose that locally optimal solution that has the least cost among all those found. As the number of kinds of local transformations that have been used to solve various problems is great, we shall close the section with two examples — the TSP, and a simple problem of "package placement."

### **The Traveling Salesman Problem**

The TSP is one for which local search techniques have been remarkably successful. The simplest transformation that has been used is called "2-opting." It consists of taking any two edges, such as (A, B) and (C, D) in Fig. 10.25, removing them, and reconnecting their endpoints to form a new tour. In Fig. 10.25, the new tour runs from B, clockwise to C, then along the edge (C, A), counterclockwise from A to D, and finally along the edge (D, B). If the sum of the lengths of (A, C) and (B, D) is less than the sum of the lengths of (A, B) and (C, D), then we have an improved tour.<sup>†</sup> Note that we cannot

 $^\dagger$  Do not be fooled by the picture of Fig. 10.25. True, if lengths of edges are distances in the plane, then the dashed edges in Fig. 10.25 must be longer than those they replace. In



Fig. 10.24. Local search in the space of solutions.

connect A to D and B to C, as the result would not be a tour, but two disjoint cycles.

To find a locally optimal tour, we start with a random tour, and consider all pairs of nonadjacent edges, such as (A, B) and (C, D) in Fig. 10.25. If the tour can be improved by replacing these edges with (A, C) and (B, D), do so, and continue considering pairs of edges that have not been considered before. Note that the introduced edges (A, C) and (B, D) must each be paired with all the other edges of the tour, as additional improvements could result.

**Example 10.12.** Reconsider Fig. 10.21, and suppose we start with the tour of Fig. 10.26(a). We might replace (a, e) and (c, d), with a total cost of 12, by (a, d) and (c, e), with a total cost of 10, as shown in Fig. 10.26(b). Then we might replace (a, b) and (c, e) by (a, c) and (b, e), giving the optimal tour shown in Fig. 10.26(c). One can check that no pair of edges can be removed from Fig. 10.26(c) and be profitably replaced by crossing edges with the same endpoints. As one case in point, (b, c) and (d, e) together have the relatively high cost of 10. But (c, e) and (b, d) are worse, costing 14 together.  $\Box$ 

general, however, there is no reason to assume the distances in Fig. 10.25 are distances in the plane, or if they are, it could have been (A, B) and (C, D) that crossed, not (A, C) and (B, D).







Fig. 10.26. Optimizing a TSP instance by 2-opting.

We can generalize 2-opting to k-opting for any constant k, where we remove up to k edges and reconnect the remaining pieces in any order so that result is a tour. Note that we do not require the removed edges to be nonadjacent in general, although for the 2-opting case there was no point in considering the removal of two adjacent edges. Also note that for k>2, there is more than one way to connect the pieces. For example, Fig. 10.27 shows the general process of 3-opting using any of the following eight sets of edges.

1. $(A, F), (D, E), (B, C)$ (as the tour wa	as)
2. $(A, F), (C, E), (D, B)$ (a 2-opt)	
3. $(A, E), (F, D), (B, C)$ (another 2-opt)	)
4. $(A, E), (F, C), (B, D)$ (a true 3-opt)	
5. $(A, D), (C, E), (B, F)$ (another true 3)	3-opt)
6. $(A, D), (C, F), (B, E)$ (another true 3)	3-opt)
7. $(A, C), (D, E), (B, F)$ (a 2-opt)	
8. $(A, C), (D, F), (B, E)$ (a 3-opt)	



Fig. 10.27. Pieces of a tour after removing three edges.

It is easy to check that, for fixed k, the number of different k-opting transformations we need to consider if there are n vertices is  $O(n^k)$ . For example, the exact number is n(n-3)/2 for k = 2. The time taken to obtain a locally optimal tour may be considerably higher than this, however, since we could make many local transformations before reaching a locally optimum tour, and each improving transformation introduces new edges that may participate in later transformations that improve the tour still further. Lin and Kernighan [1973] have found that variable-depth-opting is in practice a very powerful method and has a good chance of getting the optimum tour on 40-100 city problems.

## **Package Placement**

The one-dimensional package placement problem can be stated as follows. We have an undirected graph, whose vertices we call "packages." The edges are labeled by "weights," and the weight w(a, b) of edge (a, b) is the number of "wires" between packages a and b. The problem is to order the vertices  $p_1, p_2, \ldots, p_n$ , such that the sum of  $|i-j| w(p_i, p_j)$  over all pairs i and j is minimized. That is, we want to minimize the sum of the lengths of the wires.

The package placement problem has had a number of applications. For

example, the "packages" could be logic cards on a rack, and the weight of an interconnection between cards is the number of wires connecting them. A similar problem comes up when we try to design integrated circuits from arrays of standard modules and interconnections between them. A generalization of the one-dimensional package placement problem allows placement of "packages," which have height and width, in a two-dimensional region, while minimizing the sum of the lengths of the wires between packages. This problem also has application to the design of integrated circuits, among other areas.

There are a number of local transformations we could use to find local optima for instances of the one-dimensional package placement problem. Here are several.

- 1. Interchange adjacent packages  $p_i$  and  $p_{i+1}$  if the resulting order is less costly. Let L(j) be the sum of the weights of the edges extending to the left of  $p_j$ , i.e.,  $\sum_{k=1}^{j-1} w(p_k, p_j)$ . Similarly, let R(j) be  $\sum_{k=j+1}^{n} w(p_k, p_j)$ . Improvement results if  $L(i) R(i) + R(i+1) L(i+1) + 2w(p_i, p_{i+1})$  is negative. The reader should verify this formula by computing the costs before and after the interchange and taking the difference.
- 2. Take a package  $p_i$  and insert it between  $p_j$  and  $p_{j+1}$  for some i and j.
- 3. Interchange any two packages  $p_i$  and  $p_j$ .

**Example 10.13.** Suppose we take the graph of Fig. 10.21 to represent a package placement instance. We shall restrict ourselves to the simple transformation set (1). An initial placement, a, b, c, d, e, is shown in Fig. 10.28(a); it has a cost of 97. Note that the cost function weights edges by their distance, so (a, e) contributes  $4 \times 7 = 28$  to the cost of 97. Let us consider interchanging d with e. We have L(d) = 13, R(d) = 6, L(e) = 24, and R(e) = 0. Thus L(d)-R(d)+R(e)-L(e)+2w(d, e) = -5, and we can interchange d and e to improve the placement to (a, b, c, e, d), with a cost of 92 as shown in Fig. 10.28(b).

In Fig. 10.28(b), we can interchange c with e profitably, producing Fig. 10.28(c), whose placement (a, b, e, c, d) has a cost of 91. Fig. 10.28(c) is locally optimal for set of transformations (1). It is not globally optimal; (a, c, e, d, b) has a cost of 84.  $\Box$ 

As with the TSP, we cannot estimate closely the time it takes to reach a local optimum. We can only observe that for set of transformations (1), there are only n-1 transformations to consider. Further, if we compute L(i) and R(i) once, we only have to change them when  $p_i$  is interchanged with  $p_{i-1}$  or  $p_{i+1}$ . Moreover, the recalculation is easy. If  $p_i$  and  $p_{i+1}$  are interchanged, for example, then the new L(i) and R(i) are, respectively,  $L(i+1) - w(p_i, p_{i+1})$  and  $R(i+1) + w(p_i, p_{i+1})$ . Thus O(n) time suffices to test for an improving transformation and to recalculate the L(i)'s and R(i)'s. We also need only O(n) time to initialize the L(i)'s and R(i)'s, if we use the recurrence





# L(1)=0

$$L(i) = L(i-1) + w(p_{i-1}, p_i)$$

and a similar recurrence for R.

In comparison, the sets of transformations (2) and (3) each have  $O(n^2)$  members. It will therefore take  $O(n^2)$  time just to confirm that we have a locally optimal solution. However, as for set (1), we cannot closely bound the total time taken when a sequence of improvements are made, since each improvement can create additional opportunities for improvement.  $\Box$ 

## Exercises

- 10.1 How many moves do the algorithms for moving *n* disks in the Towers of Hanoi problem take?
- \*10.2 Prove that the recursive (divide and conquer) algorithm for the Tower of Hanoi and the simple nonrecursive algorithm described at the beginning of Section 10.1 perform the same steps.
- **10.3** Show the actions of the divide-and-conquer integer multiplication algorithm of Fig. 10.3 when multiplying 1011 by 1101.
- \*10.4 Generalize the tennis tournament construction of Section 10.1 to tournaments where the number of players is not a power of two. *Hint*. If the number of players n is odd, then one player must receive a bye

(not play) on any given day, and it will take n days, rather than n-1 to complete the tournament. However, if there are two groups with an odd number of players, then the players receiving the bye from each group may actually play each other.

10.5 We can recursively define the number of combinations of m things out of n, denoted  $\binom{n}{m}$ , for  $n \ge 1$  and  $0 \le m \le n$ , by

$$\binom{n}{m} = 1$$
 if  $m = 0$  or  $m = n$   
 $\binom{n}{m} = \binom{n-1}{m} + \binom{n-1}{m-1}$  if  $0 < m$ 

- a) Give a recursive function to compute  $\binom{n}{m}$ .
- b) What is its worst-case running time as a function of n?
- c) Give a dynamic programming algorithm to compute  $\binom{n}{m}$ . Hint. The algorithm constructs a table generally known as Pascal's triangle.

< n

- d) What is the running time of your answer to (c) as a function of n.
- **10.6** One way to compute the number of combinations of m things out of n is to calculate  $(n)(n-1)(n-2)\cdots(n-m+1)/(1)(2)\cdots(m)$ .
  - a) What is the worst case running time of this algorithm as a function of n?
  - \*b) Is it possible to compute the "World Series Odds" function P(i, j) from Section 10.2 in a similar manner? How fast can you perform this computation?
- 10.7 a) Rewrite the odds calculation of Fig. 10.7 to take into account the fact that the first team has a probability p of winning any given game.
  - b) If the Dodgers have won one game and the Yankees two, but the Dodgers have a .6 probability of winning any given game, who is more likely to win the World Series?
- 10.8 The odds calculation of Fig. 10.7 requires  $O(n^2)$  space. Rewrite the algorithm to use only O(n) space.
- \*10.9 Prove that Equation (10.4) results in exactly  $2\binom{i+j}{i} 1$  calls to P.
- 10.10 Find a minimal triangulation for a regular octagon, assuming distances are Euclidean.
- 10.11 The paragraphing problem, in a very simple form, can be stated as follows: We are given a sequence of words  $w_1, w_2, \ldots, w_k$  of lengths  $l_1, l_2, \ldots, l_k$ , which we wish to break into lines of length L. Words are separated by blanks whose ideal width is b, but blanks can stretch or shrink if necessary (but without overlapping words), so that a line  $w_i \ w_{i+1} \ \cdots \ w_j$  has length exactly L. However, the penalty

## EXERCISES

for stretching or shrinking is the magnitude of the total amount by which blanks are stretched or shrunk. That is, the cost of setting line  $w_i w_{i+1} \cdots w_j$  for j > i is (j-i) |b'-b|, where b', the actual width of the blanks, is  $(L-l_i-l_{i+1}-\cdots-l_j)/(j-i)$ . However, if j=k (we have the last line), the cost is zero unless b' < b, since we do not have to stretch the last line. Give a dynamic programming algorithm to find a least-cost separation of  $w_1, w_2, \ldots, w_k$  into lines of length L. Hint. For  $i=k, k-1, \ldots, 1$ , compute the least cost of setting  $w_i, w_{i+1}, \ldots, w_k$ .

10.12 Suppose we are given *n* elements  $x_1, x_2, \ldots, x_n$  related by a linear order  $x_1 < x_2 < \cdots < x_n$ , and that we wish to arrange these elements *m* in a binary search tree. Suppose that  $p_i$  is the probability that a request to find an element will be for  $x_i$ . Then for any given binary search tree, the average cost of a lookup is  $\sum_{i=1}^{n} p_i(d_i + 1)$ , where  $d_i$  is the depth of the node holding  $x_i$ . Given the  $p_i$ 's, and assuming the  $x_i$ 's never change, we can find a binary search tree that

assuming the  $x_i$ 's never change, we can find a binary search tree that minimizes the lookup cost. Find a dynamic programming algorithm to do so. What is the running time of your algorithm? *Hint*. Compute for all *i* and *j* the optimal lookup cost among all trees containing only  $x_i, x_{i+1}, \ldots, x_{i+j-1}$ , that is, the *j* elements beginning with  $x_i$ .

- **\*\*10.13** For what values of coins does the greedy change-making algorithm of Section 10.3 produce an optimal solution?
  - **10.14** a) Write the recursive triangulation algorithm discussed in Section 10.2.
    - b) Show that the recursive algorithm results in exactly  $3^{s-4}$  calls on nontrivial problems when started on a problem of size  $s \ge 4$ .
  - **10.15** Describe a greedy algorithm for
    - a) The one-dimensional package placement problem.
    - b) The paragraphing problem (Exercise 10.11).

Give an example where your algorithm does not produce an optimal answer, or show that no such example exists.

- 10.16 Give a nonrecursive version of the tree search algorithm of Fig. 10.17.
- 10.17 Consider a game tree in which there are six marbles, and players 1 and 2 take turns picking from one to three marbles. The player who takes the last marble *loses* the game.
  - a) Draw the complete game tree for this game.
  - b) If the game tree were searched using the alpha-beta pruning technique, and nodes representing configurations with the smallest number of marbles are searched first, which nodes are pruned?

# c) Who wins the game if both play their best?

- \*10.18 Develop a branch and bound algorithm for the TSP based on the idea that we shall begin a tour at vertex 1, and at each level, branch based on what node comes next in the tour (rather than on whether a particular edge is chosen as in Fig. 10.22). What is an appropriate lower bound estimator for configurations, which are lists of vertices 1,  $v_1$ ,  $v_2$ , ... that begin a tour? How does your algorithm behave on Fig. 10.21, assuming *a* is vertex 1?
- \*10.19 A possible local search algorithm for the paragraphing problem is to allow local transformations that move the first word of one line to the previous line or the last word of a line to the line following. Is this algorithm locally optimal, in the sense that every locally optimal solution is a globally optimal solution?
- **10.20** If our local transformations consist of 2-opts only, are there any locally optimal tours in Fig. 10.21 that are not globally optimal?

## **Bibliographic Notes**

There are many important examples of divide-and-conquer algorithms including the  $O(n\log n)$  Fast Fourier Transform of Cooley and Tukey [1965], the  $O(n\log n\log \log n)$  integer multiplication algorithm of Schonhage and Strassen [1971], and the  $O(n^{2.81})$  matrix multiplication algorithm of Strassen [1969]. The  $O(n^{1.59})$  integer multiplication algorithm is from Karatsuba and Ofman [1962]. Moenck and Borodin [1972] develop several efficient divide-andconquer algorithms for modular arithmetic and polynomial interpolation and evaluation.

Dynamic programming was popularized by Bellman [1957]. The application of dynamic programming to triangulation is due to Fuchs, Kedem, and Uselton [1977]. Exercise 10.11 is from Knuth [1981]. Knuth [1971] contains a solution to the optimal binary search tree problem in Exercise 10.12.

Lin and Kernighan [1973] describe an effective heuristic for the traveling salesman problem.

See Aho, Hopcroft, and Ullman [1974] and Garey and Johnson [1979] for a discussion of NP-complete and other computationally difficult problems.